**APPENDIX 1** 

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Robust Summaries for 1-Decene, Tetramer, Mixed with 1-Decene Trimer, Hydrogenated, and Structurally Analogous PAOs (Revised October 4, 2002)

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# **BIODEGRADATION (CAS NO.: 68649-12-7)**

Test Substance:	CAS No.: 68649-12-7; 1-Decene, tetramer, mixed with 1-decene trimer, hydrogenated
Method/Guideline:	USEPA EPA 560/6-83-003, CG-2000
Year (guideline):	1982
Type (test type):	Aerobic Aquatic Biodegradation
GLP:	Yes
Year (study performed):	1992
Inoculum:	Domestic activated sludge
Exposure Period:	28 days
Note: Concentration prep., vessel type, replication, test conditions.	Activated sludge and test medium were combined prior to test material addition. The media consisted of a mineral salt solution, activated sludge, and distilled water.  The test system utilized 2.0L Erlenmeyer flasks as test vessels. The test and reference materials were added to duplicate test vessels followed by enough volume of test medium to yield a 1.0 L final volume, after inoculum addition. Mixed liquor was added to each flask to give a final dry sludge solids concentration of 30 mg/L. In addition, 0.1 g of soil was added to each flask followed by 1 mL of yeast extract solution (15 mg yeast extract per 100 mL of distilled water). The flasks were closed with neoprene stoppers containing a 10 mL KOH trap and an inlet and outlet port. The flasks were then placed on a rotary shaker at 25°± 3°C at approximately 150 rpm.  Potential biodegradability was evaluated at two test material loadings of approximately 10 mg/L and 20 mg/L of carbon. Sodium benzoate (positive control) concentration was approximately 20 mg/L of carbon.  Twice per week, the flasks were monitored for spent NaOH and evolved carbon dioxide (CO <sub>2</sub> ) was determined by titration.
Results: Units/Value:	On day 28, 53.5 and 49.2% biodegradation was achieved in the 10 and 20 mg/L carbon loadings, respectively.
Note: Deviations from protocol or guideline, analytical method.	The positive control (sodium benzoate) degraded by 84% by day 28.  No excursions from the protocol were noted.
Conclusion:	The test material did not meet the criteria to be classified as readily biodegradable, however, the test material did biodegrade to

	a great extent which suggests that it will not persist in the environment.
Reliability:	(1) Reliable without restriction
Reference:	Mobil Oil Corporation, Environmental and Health Sciences Laboratory. 1992. Aerobic Biodegradation Study, SHF-41, Study No. 65226.
Other (source):	ExxonMobil Biomedical Sciences, Inc.

# FISH ACUTE TOXICITY (CAS NO.: 68649-12-7)

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Test Substance:	CAS No.: 68649-12-7; 1-Decene, tetramer, mixed with 1-decene trimer, hydrogenated	
Method/Guideline:	Fish Acute Toxicity Test (EPA 560/6-82-002; OECD 203)	
Type (test type):	Fish Acute Toxicity Test; Dispersion Test	
GLP:	Yes	
Year (study performed):	1992	
Species:	Sheepshead Minnow (Cyprinodon variegatus)	
Analytical Monitoring:	No	
Exposure Period:	96 hour	
Statistical Method:	Binomial Probability Analysis (Stephan, et al., 1978)	
Note: Concentration prep. vessel type, volume, replication, water quality parameters, environmental conditions, organisms supplier, age, size, weight, loading.	Individual Oil/Water Dispersion (OWD) systems were prepared for each treatment level. The test system was designed to maintain the test substance as a dispersion of small droplets throughout the water column. The test substance was added volumetrically, via graduated cylinder, directly to the dispersion system. Each test chamber was a 10-gallon glass aquaria with 30 liters of water, and was equipped with a vertically mounted, motor-driven impeller assembly. The impeller assembly, consisting of 3-blades on a 10-inch stainless steel shaft, was housed in a 2-inch diameter PVC cylinder with 4 horizontal apertures near the bottom. Water and test substance spilling into the top of the cylinder were expelled through the apertures at the bottom. The OWD systems operated continuously for the duration of the test. Twenty fish were randomly assigned to each chamber. No renewal of the test solutions was performed during the test.  Test temperature was 19.5 - 20.0 Deg C., lighting was 16 hours light: 8 hours dark. Dissolved oxygen ranged from 6.7 to 7.0 mg/L, and pH ranged from 8.1 to 8.4 during the study. Salinity ranged from 18.4 - 21.9 ppt. Fish were not fed during the study.	

	Fish Mean Wt.(Control) = 0.55g; mean standard length = 2.9 test organism loading = 0.37 g of fish/L.	9cm,
Results: Units/Value:	96-hr LL0 = 5002mg/L, based on nominal loading levels.	
Note: Deviations from protocol or guideline, analytical method, biological observations, control survival.		
Results continued	Loading Level. % Mortality @ 96 hr.	
	Control 10	
	98 mg/L 0	
	492 mg/L 0	
	1011 mg/L 0	
	2023 mg/L 0	
	5002 mg/L 0	
Conclusion:		
Reliability:	Code 1, Reliable without restriction	
Reference:	Mobil Oil Corporation, Environmental and Health Sciences Laboratory. 1992. Static 96-hour Acute Toxicity Study of SHF-41 to Sheepshead Minnow, Study No. 65227.	
Other (source):	ExxonMobil Biomedical Sciences, Inc.	

# **INVERTEBRATE ACUTE TOXICITY (CAS NO.: 68649-12-7)**

Test Substance:	CAS No.: 68649-12-7; 1-Decene, tetramer, mixed with 1-decene trimer, hydrogenated
Method/Guideline:	Mysid Acute Toxicity Test (EPA 560/6-82-002)
Type (test type):	Mysid Acute Toxicity Test - Static Renewal
GLP:	Yes
Year (study performed):	1992

Species:	Mysid Shrimp ( <i>Mysidopsis bahia</i> )
Analytical Monitoring:	No
Exposure Period:	96 hour
Statistical Method:	Binomial Probability Analysis (Stephan, et al., 1978)
Note: Concentration prep. vessel type, volume, replication, water quality parameters, environmental conditions, organisms supplier, age, size, weight, loading.	Individual Water Accomodated Fractions (WAF) were prepared for each treatment solution twenty-four hours prior to test initiation. WAFs were stirred with magnetic stir bars and stirplates in glass aspirator bottles containing 1 L of test water. A measured amount of the test material was pipetted onto the water surface of each bottle. All aspirator bottles were covered with parafilm to minimize evaporation. Stirring produced a vortex of less than 25% of the depth of the solution. After the stirring/settling period, the aqueous phase (WAF) was drawn from the outlet at the bottom of each aspirator bottle and split into two replicates of 400 mL each. This procedure was repeated to prepare fresh solutions at 24, 48, and 72 hours.  Test chambers were 1 L Pyrex crystallizing dishes. Ten mysids were randomly assigned to each chamber.  Test temperature was 19.8 - 20.6 Deg C; lighting was 16 hours light: 8 hours dark. Dissolved oxygen ranged from 6.2 to 7.2 mg/L and pH was approximately 8.4 during the study. Salinity ranged from 19.0 - 22.0 ppt. Mysids were fed newly hatched <i>Artemia</i> spp. nauplii (approximately 24 hours old) <i>ad libitum</i> prior to and daily during the test. Mysids were four to six days old at study initiation.
Results: Units/Value:	96-hr LL0 = 5002mg/L, based on nominal loading levels.
Note: Deviations from protocol or guideline, analytical method, biological observations, control survival.	
Results continued	Loading Level % Mortality @ 96 hr.
	Control 5
	82 mg/L 5
	574 mg/L 15
	1066 mg/L 5
	2050 mg/L 10
	5002 mg/L 0

Conclusion:	
Reliability:	Code 1, Reliable without restriction
Reference:	Mobil Oil Corporation, Environmental and Health Sciences Laboratory. 1992. Static-Renewal 96-hour Acute Toxicity Study of the Water Soluble Fraction (WSF) of SHF-41 to <i>Mysidopsis bahia</i> , Study No. 65228.
Other (source):	ExxonMobil Biomedical Sciences, Inc.

<b>Acute Inhalation Toxic</b>	ity (CAS NO.68037-01-4)
Test Substance CAS No.	1-Decene Homopolymer, hydrogenated 68037-01-4
Method/Guideline Type of Study GLP Year Species/strain Sex No. of animals/sex/dose Route of administration Dose/Concentration Levels Exposure Time	Other Acute Inhalation Yes 1988 Sprague-Dawley rats Males/Females 10/sex Inhalation - Aerosol 0, 0.48, and 2.5 mg/L 4 hours
Remarks on Test Conditions	The test atmosphere was generated with a Laskin nebulizer to aerosolize the test material. The concentration of the aerosol was controlled by varying the design of the nebulizer, the air pressure, and air flow. A glass elutriator was used to remove large particles from the aerosol to maximize the percentage of respirable particles in the aerosol stream. The aerosol concentration in the chamber was determined gravimetrically.
Results	Young adult rats were exposed for 4 hours to 0, 0.48, or 2.5 mg/L of aerosolized test material for 4 hours. Half of the animals in each group were sacrificed the day after exposure and half were sacrificed two weeks after exposure. On the day of sacrifice, animals were necropsied, wet and dry weights of kidneys, liver, and right middle lung lobe were measured, and selected tissues were preserved for histopathology. From the control and high dose groups, the following tissues were examined microscopically: nasal turbinates, tracheobronchial lymph nodes, left kidney, liver, and all gross lesions.
Remarks	$LC_{50} > 2.5 \text{ mg/L}$
	The mean aerosol concentrations for the exposed groups were 0.48 and 2.5 mg/L. The mass median aerodynamic diameter of the generated particles was 1.1 $\mu$ m. All animals appeared normal

during the exposure, although less than one-half of the animals in the high-dose group could be observed because of poor visibility within the test chamber due to the high aerosol concentration. No animals died during the exposure, or during the following two-week observation period. No toxicologically significant changes were observed for clinical signs and mean body weights of the animals following exposure. Organ weights were not affected by exposure. The lung was the only organ with treatment-related changes. At the 1-day sacrifice, dark depressed or discolored areas were observed in 3 animals from the middle dose group and 6 animals from the high dose group. However, since none of these lesions were found at the 2-week sacrifice, the effects appeared to be reversible. Acute focal inflammation was observed in 2 animals from the 0.5 mg/L group and 6 animals from the 2.5 mg/L group. These foci were present in the respiratory tract (along the route of exposure) and resolved by 2 weeks following the exposure.

Conclusions

Under the conditions of this study, decene homopolymer has a low order of acute toxicity via the inhalation route of exposure.

**Data Quality** 

1 - Reliable without restrictions.

Reference

Final Report on Acute Inhalation Toxicity of Decene Homopolymer, (1987) Mobil Environmental and Health Science Laboratory

Date last changed

December 2001

## Repeated Dose Toxicity (CAS No.: 68037-01-4)

Test Substance 1-Decene Homopolymer, hydrogenated

CAS No. 68037-01-4

Method/Guideline Other

Type of Study Subchronic Range-finding Study

GLP Yes Year 1990

Species/strain Sprague Dawley

Sex Females
No. of animals/sex/dose 5/dose
Route of administration Oral gayage

Frequency of treatment 5 days per week for 4 weeks Dose/Concentration Levels 0, 500, 2500, 5000 mg/kg/day

Control group and treatment 2 Control groups: Sham treatment and untreated

Remarks on Test Conditions Female rats (5/dose), in addition to a sham-gavage control group and

an untreated control group, were individually housed and allowed free access to food and water. Animals were checked for morbidity and mortality at least once daily. Animals were weighed immediately before the first dosing and approximately weekly thereafter.

Macroscopic findings were noted at necropsy, but organ weights were not measured. The liver was fixed and examined microscopically.

Results NOAEL = 5000 mg/kg/day

Remarks

No deaths occurred during the study. There were no significant changes in body weight. The only clinical signs that could be attributed to the test material were oily staining around the anus and soft stool. No gross pathological changes were observed in any of the groups. Histological evaluation of the liver revealed no adverse effects. The results of this study were used to set the doses for a 90-day feeding study.

Conclusions

Under the conditions of this study, decene homopolymer has a low

order of subchronic toxicity via the oral route of administration.

Reliability 2 - Reliable with restrictions - rangefinding study Reference Range-Finding Study: Oral Administration of Una

Range-Finding Study: Oral Administration of Unadditized decene homopolymer to rats (1990), Performed by Mobil Environmental

Health and Safety Department.

Date last changed December, 01

## REPEATED DOSE TOXICITY (CAS NO.: 68037-01-4)

Test Substance CAS No.	1-Decene Homopolymer, hydrogenated 68037-01-4
Method/Guideline  Type of Study GLP Year Species/strain Sex No. of animals/sex/dose Route of administration Frequency of treatment Dose/Concentration Levels Control group and treatment	FDA Guideline: Guidelines for Subchronic Oral Toxicity Studies EPA Guideline: Health Effects Test Guidelines: Subchronic Exposure 90-Day Oral Administration in Diet Yes 1990 Sprague-Dawley Rats Males/Females 20/sex/dose Diet Ad libitum, 13 weeks 20,000 ppm in diet Control Diet
Remarks on Test Conditions	Rats (20/sex/dose) were fed lab chow containing 0, 500, 5000, and 20,000 ppm decene homopolymer for 90 days. Diet was prepared one week prior to initiation of the study and every other week thereafter. Feed homogeneity and stability analyses were performed on each batch. Animals had continuous access to test diet and water. The amount of food consumed was determined three times per week. Each animal was observed daily during the course of the study for clinical signs, mortality, and moribundity. Blood samples were collected prior to treatment and both blood and urine samples were collected during weeks 5 and 13. At the end of the exposure period, animals received ophthalmologic examinations and were euthanized and necropsied with a complete gross examination. Fresh organ weight was determined in 10 males and 9 females. From each rat, 50 tissues were preserved. From these, tissues of the control and high dose groups were examined by light microscopy. Gross pathology data were evaluated statistically using by ANOVA.
Results	

NOAEL = 20,000 ppmRemarks All rats survived to study termination. No clinical signs indicative of systemic toxicity were observed during this study. The test material did not adversely effect body weight gain or food consumption. Urinalysis and ophthalmologic exam data did not indicate any treatment-related effects. No statistically significant changes were observed in the hematology analysis between the control and treated animals. Statistically significant differences were found between the serum chemistry data from the untreated and treated animals. Specifically, a linear relationship was found between the dose and serum level for albumin/globulin ratio in males and for inorganic phosphorous in females. When compared to historical serum reference values, only the dose-response curve for the inorganic phosphorous at the highest dose fell outside the normal range of the historical data. Animals in the study exhibited no signs of compound related ocular disease. Urinalysis did not reveal any treatment-related changes. There were no changes in organ weight that were deemed to be due to treatment. No effects on the enteric tract were observed. Mean liver weight and microscopic examinations of the liver did not vary significantly between the high dose and control rats. None of the major organs or organ systems, including male and female reproductive organs, showed any detectable treatment-related changes. Conclusions Decene homopolymer did not produce any toxicologically significant effects under the conditions of this study. Reliability 1 - Reliable without restrictions Reference 90-Day Oral Administration of Unadditized Decene Homopolymer in the Diet of Rats. (1990) Performed by Mobil Environmental Health and Safety Department.

## REPEATED DOSE TOXICITY (CAS NO.: 68037-01-4)

December, 01

Date last changed

Test Substance 1-Decene Homopolymer, hydrogenated CAS No. 68037-01-4 Method/Guideline **OECD 408** Type of Study 90-Day Subchronic GLP Yes Year 1995 Species/strain Fischer 344 Rat Sex M/F No. of animals/sex/dose 10/sex/group Route of administration Diet Dose/Concentration Levels 200, 20,000 ppm Control group and treatment Control Diet Remarks on Test Conditions Decene Homopolymer at dose levels of 200 and 20,000 ppm was administered in the diet to male and female Fischer 344 rats. Diet was prepared 1-week prior to administration and every two weeks thereafter. Food consumption was measured three times per

week. Virus-free Fischer 344 rats were received at about 4 weeks of age. Prior to treatment, blood samples were drawn and animals received an ophthalmologic examination. Following a 2-week quarantine, animals were randomly allocated to treatment groups (10/sex/group), individually housed, and received test material in the diet at 200 and 20,000 ppm. Food and water were available *ad libitum*. Animals were checked for morbidity and mortality at least once daily. Hematology and serum chemistry analyses were performed on blood samples collected prior to treatment, and during weeks 5 and 13. Ophthalmoscopic examinations of the eyelids, bulbar conjunctiva, cornea, iris, anterior chamber, and lens were performed. Necropsies were performed at the end of the study and tissues were collected for histopathologic evaluation. Results were analyzed by ANOVA and associated by F-test.

Results NOAEL > 20,000 ppm

Remarks

No clinical signs were observed during the study. Aside from two animals that died during the 13-week blood collection, all animals survived until the end of the study. In general the amount of food consumed by treated animals was comparable to controls and body weight gain was normal. Hematology results were normal in both

treated and control animals. However, there were statistically significant differences in the serum chemistry data, (glucose in males, and sodium, phosphorous, and calcium in females) between the control and treated animals. These differences were considered

marginal and the biological significance was not clear.

Ophthalmoscopic examinations revealed no abnormalities. The test material did not produce any significant findings at necropsy. In addition, there were no significant treatment-related changes in the

liver or mesenteric lymph nodes.

Conclusions Decene Homopolymer poses a low order of subchronic toxicity.

Data Quality 1 - Reliable without Restrictions

Reference 90-Day Oral Feeding Study in Fischer 344 Rats with Hydrogenated

Polyalpha Decene in the Diet, (1995) Performed by Stonybrook

Laboratories, Inc. for Mobil Corporation.

Date last changed December, 2001

## **REPEATED DOSE TOXICITY (CAS NO. 68037-01-4)**

Remarks	1-Decene Homopolymer, hydrogenated (31.3% trimer, 45.0% tetramer, 23.7% pentamer and higher), CAS# 68037-01-4
Method	
Method/guideline followed	Other
Test type	Combined repeated exposure and reproduction oral toxicity study
GLP	Yes.
Year	1994
Species	Rat

Strain

Route of administration

**Duration of test** 

Crl:CD® (Sprague-Dawley) BR VAF/Plus

Oral gavage

Parental males were dosed for 4 wks prior to mating, through 15-day mating period.

Parental females were dosed for 4 wks prior to mating, through pregnancy and until day 20 post-partum.

Offspring were dosed for 91 days starting on day 22 post-partum.

0, 100, 500, 1000 mg/kg/day

F0: 30 males, 30 females per group; F1: 20 males, 20 females per group.

7 davs/week

Polyethylene glycol 400, 5 ml/kg

Not applicable.

Doses/concentration levels Sex Frequency of treatment Control group and treatment Post exposure observation

period

Statistical methods

Adult body weights, body weight gains, feed consumption, organ weights, clinical chemistry data and appropriate hematologic data were evaluated by ANOVA. When significance was observed with ANOVA, group by group comparisons were performed using Dunnett's Test or a modified version of Dunnett's Test. All tests were two-tailed with a minimum significance level of 5% comparing the control group to each treatment

group.

**Test Conditions** 

The study design included a 91-day main study for repeated dose toxicity end points and a reproductive toxicity study (summarized separately). Animals utilized for the 91-day toxicity study were offspring of animals administered the test article for 4 weeks prior to mating, during mating, and, for females, through gestation and lactation day 20.

### Parental animals:

Animals were observed daily for overt signs of toxicity. Males were weighed weekly. Females were weighed weekly prior to mating and on gestation days 0,7,14, and 20 and lactation days 1,7,14, and 21. Food consumption was measured on the same days as body weights, except during cohabitation. All animals were subjected to a gross necropsy.

### Offspring:

Animals were observed daily for overt signs of toxicity. Body weights and food consumption were measured weekly. Ophthalmology examinations were performed on all animals prior to initiation of dosing and near study conclusion. Blood samples were obtained on the day of euthanasia for evaluation of clinical pathology parameters. A gross necropsy examination was performed on all animals. Organ weights were obtained for liver, kidneys, thyroid/parathyroid, adrenal glands, gonads and brain. Histopathologic examinations were performed on all control and high animals and animals found dead during the study.

The study also contained reproductive and developmental toxicity endpoints (summarized separately).

Results

NOAEL (NOEL) LOAEL (LOEL)

1000 mg/kg/day. Not applicable.

Remarks

No treatment related toxicity was observed in the F0 male and female rats. The F1 pups did not demonstrate any test article related toxicity during parturition and lactation. In the F1 rats during the 91-day toxicity phase, clinical observations representing minor gastrointestinal disturbances were seen in all groups and were judged to be vehicle related. No apparent test

article related clinical observations were noted. Transient changes in body weights, weight gain, food consumption, hematology parameters and organ weights were seen at a few intervals, but were not considered to be biologically meaningful. A statistically significant increase in prothrombin time was seen in the males of the 1000 mg/kg/day group, however, this change did not correlate with a decrease in platelets, gross necropsy findings or any lesions noted histopathologically, Therefore, this increase in prothrombin time was not considered to be biologically meaningful.

### **Conclusions**

Repeated oral exposure of 1-decene homopolymer, hydrogenated, to male and female Sprague Dawley rats at levels of 0, 100, 500, and 1000 mg/kg/day produced no evidence of any adverse effects on clinical observations, organ weights, gross or histopathology, clinical chemistry or hematology endpoints. Based on these data, the no-observable-effect level (NOEL) for repeated dose toxicity was 1000 mg/kg/day, the highest concentration tested.

<u>Data Quality</u> Reliabilities **References** 

Klimish value = 1 (Reliable without restrictions).

Daniel, E.M. (1994) An oral (gavage) 91-day toxicity study of EthylFlo 166 in rats with an <u>in utero</u> exposure phase. Report of Springborn Laboratories, Inc., Spencerville, OH, conducted for Albemarle

Corporation, Baton Rouge, LA.

<u>Other</u>

Last changed 31-October-01

Genetic Toxicity 'In Vivo' (CAS No.: 68037-01-4)

Test Substance 1-Decene Homopolymer, hydrogenated

CAS No. **68037-01-4** 

Method Other

Type of Study Micronucleus/Subchronic Dermal

GLP Yes Year 1985 Species/Strain Rats

No. of 15/sex/dose

animals/sex/dose

Remarks on Test

Conditions daily, 5 days per week for 13 weeks at doses of 0, 800, or 2000

mg/kg/day. Subchronic dermal exposure was selected as the exposure method because it is the most relevant route of exposure for industrial use. At the end of the 13-week period, appropriate tissue was harvested for micronucleus evaluation. Femurs were taken from 5 rats/sex/dose and peripheral blood smears were made. Slides were air dried, fixed in methanol and then stained with acridine orange. The incidence of micronuclei was determined per 1000 polychromatic erythrocytes

15 Rats/sex/dose were exposed dermally to Decene Homopolymer

(PCEs) or normochromatic erythrocytes (NCEs). Potential cytotoxicity of the test material was monitored and determined as the ratio of

polychromatic and normochromatic erythrocytes. Results were analyzed

by ANOVA.

Results NOAEL > 2000 mg/kg/day

Remarks for Results Decene Homopolymer was not cytotoxic to red blood cell formation as

shown by the ratio of PCEs to NCEs. No statistically significant increases in the formation of micronucleated polychromatic or normochromatic erythrocytes were observed at any dose.

Conclusions Decene Homopolymer is not clastogenic following subchronic dermal

application to rats.

**Data Quality** 2 - Reliable with restrictions - not a guideline study

Reference Micronucleus assay of bone marrow and peripheral red blood cells from

rats treated via dermal administration of Synthetic Hydrocarbon-Hydrogenated Polyolefins, (1985), Mobil Environmental and Health

Science Laboratory.

Date last changed December, 01

# TOXICITY TO REPRODUCTION (CAS NO.: 68037-01-4)

**Test Substance** Remarks 1-Decene Homopolymer, hydrogenated (31.3% trimer, 45.0% tetramer, 23.7% pentamer and higher), CAS# 68037-01-4

Method

Method/auideline followed

Test type

**GLP** Year **Species** 

Strain

**Duration of test** 

Route of administration

Doses/concentration levels

Sex

Frequency of treatment Control group and

treatment

Post exposure observation

period

Statistical methods

Other

Combined repeated exposure and reproduction oral toxicity study

Yes. 1994 Rat

Crl:CD® (Sprague-Dawley) BR VAF Plus

Oral gavage

Parental males were dosed for 4 wks prior to mating, through 15-day

mating period.

Parental females were dosed for 4 wks prior to mating, through

pregnancy and until day 20 post-partum.

Offspring were dosed for 91 days starting on day 22 post-partum.

0, 100, 500, 1000 mg/kg/day

F0: 30 males, 30 females per group; F1: 20 males, 20 females per

group.

7 days/week

Polyethylene glycol, 5 ml/kg

Not applicable.

Continuous data, including body weights, weight gain, feed consumption, pre-implantation loss, gestation length, mean live litter size, implantation scar counts, clinical pathology and organ weights, were analyzed by ANOVA. When significance was observed with ANOVA, group by group comparisons were performed using Dunnett's Test or a modified version of Dunnett's Test. Count data were analyzed using Chi-Square test for copulation, fertility and pup sex ratios, the number of live and dead pups per group on lactation day 0, and pup survival after lactation day 0. Mann-Whitney U Test was utilized for resorptions. All tests were two-tailed with a minimum

significance level of 5% comparing the control group to each treatment group.

### **Test Conditions**

The study design included a 91-day main study for repeated dose toxicity end points (summarized separately) and a reproductive toxicity study. Animals utilized for the 91-day toxicity study were offspring of animals administered the test article for 4 weeks prior to mating, during mating, and, for females, through gestation and lactation day 20.

### Parental animals:

Animals were observed daily for overt signs of toxicity. Males were weighed weekly. Females were weighed weekly prior to mating and on gestation days 0,7,14, and 20 and lactation days 1,7,14, and 21. Food consumption was measured on the same days as body weights, except during cohabitation. All animals were subjected to a gross necropsy. In females, the number of implantation scars were recorded. Mating, conception and fertility indices were evaluated.

### Offspring:

The F1 offspring were evaluated for sex, viability, growth and development during lactation and the post-weaning period.

### Results

NOAEL (NOEL) LOAEL (LOEL) 1000 mg/kg/day Not applicable.

#### Remarks

There were no treatment related effects on the parental animals or on any of the reproductive parameters evaluated in this study, at any dose level. These included measures of reproductive performance (mating, conception and fertility, time to mating, gestation length, litter size), offspring survival (gestation and postnatal survival indices, percent pre- and post-implantation loss), pup body weight and pup sex ratio. No treatment related effects on reproductive organ weight or histopathology were seen in the 91-day toxicity study with the F1 animals.

#### **Conclusions**

Repeated oral exposure of 1-decene homopolymer, hydrogenated, to male and female Sprague Dawley rats at levels of 0, 100, 500, and 1000 mg/kg/day produced no evidence of adverse effects on any measures of reproductive function. Based on these data, the no-observable-effect level (NOEL) for reproductive toxicity was 1000 mg/kg/day, the highest concentration tested.

### Data Quality

Reliabilities

Klimish value = 1 (Reliable without restrictions).

#### References

Daniel, E.M. (1994) An oral (gavage) 91-day toxicity study of EthylFlo 166 in rats with an <u>in utero</u> exposure phase. Report of Springborn Laboratories, Inc., Spencerville, OH, conducted for Albemarle Corporation, Baton Rouge, LA.

## **Other**

Last changed

31-October-01

# **DEVELOPMENTAL TOXICITY (CAS NO.: 68037-01-4)**

Test Substance	1-Decene Homopolymer, hydrogenated	
CAS No.	68037-01-4	
Method/Guideline	Other	
Type of Study	Developmental	
GLP	Yes	
Year	1988	
Species/strain	Sprague-Dawley Rats	
Sex	Female	
No. of animals/sex/dose	15 females/dose	
Route of administration	Dermal	
Vehicle	None	
Frequency of Treatment	Once daily	
Dose/Concentration Levels	0, 800, and 2000 mg/kg	
Dose/Concentration Levels	0, 600, and 2000 mg/kg	
Statistics	ANOVA, Fisher's Exact Test, F-test, Student-Newman-Keul's multiple comparison test.	
Remarks on Test Conditions	Male and female rats were housed together at a ratio of 1:1. Females were examined daily for evidence of mating. Pregnant female rats were randomly assigned to dose groups (15/dose) and were dosed once daily at 0, 800, and 2000 mg/kg during GD 0-19. The test material was applied once daily to the clipped, intact dorsal skin of the rat. The test material contained 10 ppm of an antioxidant. The dermal route of exposure was chosen since it is the most relevant route of exposure in industrial applications. Rats were fitted with Elizabethan-style collars to minimize ingestion of the test material. The collars were replaced as needed throughout the gestation period. The application sites were not covered. To ensure that the rats were dosed correctly, the amount of test material to be applied to each rat was calculated using the most recently recorded body weight, dose level, and the density of the material. Control rats were clipped and collared in the same fashion as the treated rats and stroked with the tip of a syringe. Throughout gestation, animals were monitored daily for changes in appearance, behavior, or excretory function and for any signs of mortality or morbidity. Blood samples were collected from females on GD 20. The quantity or activity of 22 serum components in collected blood samples was analyzed by a flame photometer. A gross necropsy was performed on each female and the ovaries and uterus of each rat were excised and examined grossly. The following parameters were recorded: the number of corpora lutea per ovary, uterine weight, number and location of implantations, early and late resorption, and live and dead fetuses. Each fetus was gendered, weighed, measured, and grossly examined for visceral anomalies and half of the fetuses were examined for skeletal anomalies.	
Results	NOAEL = 2000 mg/kg/day	
. Counto	1107.22 2000 Highlighay	
Remarks	Rats in the two treatment groups showed minimal if any irritation at	

	the site of application of the test substance. There were no differences in food consumption during gestation between treated and control rats. Although there was a significantly smaller weight gain during gestation interval 13-16 in the high dose group, the overall weight gain for the entire gestational period was not significantly different from the controls. Serum triglycerides and albumin showed statistically significant changes between control and treated groups, however the changes did not occur in a dose-dependent fashion. It was concluded that dermal treatment of female rats during gestation had no significant effect on normal serum chemistry. At necropsy, there were no findings attributable to exposure to the test material. Reproductive performance, in utero survival, and development of the offspring were not affected by treatment. Individual body weights and crown-rump lengths of the fetuses were not altered by treatment. External, visceral, and skeletal examinations of the fetuses did not reveal any remarkable findings.
Conclusions	Under the conditions of this study, Decene Homopolymer does not induce developmental toxicity in rodents following dermal
Data Quality	application.  1 - Reliable without restrictions
Reference	Developmental Toxicity Screen in Rats Exposed Dermally to Decene
	Homopolymer, (1988), Mobil Environmental and Health Sciences Laboratory.
Date last changed	December, 01

# CARCINOGENICITY (CAS NO.: 68037-01-4)

Test Substance	1-Decene Homopolymer, hydrogenated
CAS No.	68037-01-4
Method/Guideline	Other
Type of Study	Dermal Carcinogenicity
GLP	Yes
Year	1990
Species/strain	C3H mice
Sex	Males
No. of animals/sex/dose	50 mice
Route of administration	Dermal
Frequency of Treatment	2 times per week, 104 weeks
Vehicle	None
Remarks on Test Conditions	Decene homopolymer was applied at a dose of 50 µl/application to the interscapular skin twice weekly for 104 weeks. Repeated dermal exposure is the primary route of exposure with industrial use. Mice were examined daily for skin growths, each suspected tumor was collected at the death of the mouse and examined microscopically. Male mice were received at 5-6 weeks of age and quarantined for two weeks. Mice were randomly assigned to treatment groups. During the

	study, mice were shaved approximately once a week or when necessary. Food and water were available ad libitum. Animals were observed twice daily for morbidity and moribundity. Any animals found moribund or that appeared moribund from the weight and size of the tumor or old age were sacrificed.
Results	Negative
Remarks	No treatment-related tumors were seen. In the negative control group, no primary skin tumors developed. Changes to the skin were minimal and nonspecific. Some hyperplasia was observed and deemed to be due to repeated hair removal. In the positive controls, 47 of the 50 mice developed skin tumors. Squamous cell carcinoma was the most common tumor. All mice in this group died by the 37th week. Survival of the treated group (56%) was greater than in the untreated group (42%). In the treated group, skin changes were similar to the negative controls. The interscapular skin did not have any tumors.
Conclusions	Under the conditions of this study, Decene Homopolymer is not carcinogenic in mice following chronic dermal exposure.
Data Quality	1 - Reliable without restrictions
Reference	Dermal Carcinogenicity Study on Decene Homopolymer(1990), Performed by Kettering Laboratory for Mobil Environmental and Health Science Laboratory.
Date last changed	December, 01

# **ACUTE ORAL TOXICITY (CAS NO.: 151006-60-9)**

Test substance 1-decene/1-dodecene copolymer, hydrogenated (Alkane 5)

prepared from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer (C50) and higher] and C12 oligomers. CAS 151006-

60-9

Type LD50 Species rat

StrainSprague-DawleySexmale/female

Number of animals 10

**Vehicle** other: none **Value** > 5000 mg/kg bw

Method OECD Guide-line 401 "Acute Oral Toxicity"

Year 1995 GLP Yes

### **Test condition**

A study was performed to assess the acute oral toxicity of the test material in the Sprague-Dawley strain rat. Following a range-finding study, a group of ten fasted animals (five males and five females) was given a single oral dose of undiluted test material at a dose level of 5000 mg/kg bodyweight. Individual bodyweights were recorded on the day of dosing to allow calculation of individual treatment volumes and on Day 14. The animals were observed for deaths or overt signs of toxicity 1, 2.5, and 4 hours after dosing and subsequently once daily for 14 days. They were then killed and subjected to a gross necropsy.

Result

There were no deaths. No signs of systemic toxicity were noted during the study. All animals showed expected gain in body weight during the study. No abnormalities were noted at necropsy. The acute oral median lethal dose (LD50) of the test material in the Sprague-Dawley strain rat was found to be greater than 5000 mg/kg bodyweight.

Conclusion

The acute oral median lethal dose (LD50) of the test material in the Sprague-Dawley strain rat was found to be greater than 5000 mg/kg

bodyweight.

Reliability

(1) valid without restriction

Reference

SafePharm Laboratories Limited (1995). Acute Oral Toxicity Study in The Rat. Conducted for Chevron Research and Technology Company, unpublished report.

13.12.2001

### **ACUTE INHALATION TOXICITY (CAS NO.: 151006-60-9)**

Test substance 1-decene/1-dodecene copolymer, hydrogenated (Alkane 5) prepared

from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer

(C50) and higher] and C12 oligomers. CAS 151006-60-9

Type LC50 Species rat

Strain Sprague-Dawley Sex male/female

Number of animals 10

Vehicleother: noneExposure time4 hour(s)Value> 5.06 mg/l

Method OECD Guide-line 403 "Acute Inhalation Toxicity"

Year 1995 GLP Yes

### **Test Condition**

A study was performed to assess the acute inhalation toxicity of the test material, as supplied, by exposing a single group of Sprague-Dawley strain rats (five males and five females) to an aerosol atmosphere. The animals were exposed for four hours using a nose only exposure system.

Prior to the start of the study, test material atmospheres were generated within the exposure chamber. During these periods air flow settings, test material input and the sampling system were varied to achieve the required atmospheric concentrations. During the exposure period, temperature, relative humidity, oxygen concentrations and nominal atmospheric concentrations were monitored at regular intervals. The particle size of the generated atmosphere of the test material inside the exposure chamber was determined four times during the exposure period using a Cascade Impactor.

Clinical observations were performed hourly during the exposure, immediately at the end of the exposure, one hour after the termination of the exposure and once daily for 14 days. Individual bodyweights were recorded on the day of exposure and on Days 7 and 14. Necropsies were performed on all animals at study termination.

#### Result

The mean achieved atmosphere concentration was 5.0 mg/L. The mean mass median aerodynamic diameter was 1.3 u. The inspirable fraction (%<4 u) was 90.0%. The geometric standard deviation was 0.42u. No deaths occurred. Common abnormalities noted during the study were wet fur, hunched posture, and piloerection. Incidents of decreased and increased respiratory rate, ptosis and red/brown staining around the eyes were noted. All animals appeared normal 2-3 days following exposure and for the remainder of the study. Normal bodyweight gain was noted during the study. One animal showed dark patches on the lungs but otherwise no abnormalities were detected at necropsy. The acute inhalation median lethal concentration (LC50) of the test material, Alkane 4, in the Sprague-Dawley strain rat was greater than 5.0 mg/L.

### Conclusion

The acute inhalation median lethal concentration (LC50) of the test material, Alkane 5, in the Sprague-Dawley strain rat was greater than 5.0 mg/L.

#### Reliability

(1) valid without restriction

### Reference

SafePharm Laboratories Limited (1995). Acute Inhalation Toxicity Study (Nose Only) in The Rat. Conducted for Chevron Research and Technology Company, unpublished report.

13.12.2001

## **ACUTE DERMAL TOXICITY (CAS NO.: 151006-60-9)**

### **Test substance**

1-decene/1-dodecene copolymer, hydrogenated (Alkane 5) prepared from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer (C50) and higher] and C12 oligomers. CAS 151006-60-9

Type LD50 Species rat

Strain Sprague-Dawley
Sex male/female

Number of animals 10

Vehicle other: none value > 2000 mg/kg bw

Method OECD Guide-line 402 "Acute dermal Toxicity"

Year 1995 GLP Yes

**Test condition** A study was performed to assess the acute dermal toxicity of the test

material in the Sprague-Dawley strain rat. A group of ten animals (five males and five females) was given a single, 24-hour, semi-occluded,

dermal application to intact skin at a dose level of 2000 mg/kg

bodyweight. Individual bodyweights were recorded on the day of dosing to allow calculation of individual treatment volumes and on Days 7 and 14. The animals were observed for deaths or overt signs of toxicity 1, 2.5, and 4 hours after dosing and subsequently once daily for 14 days. The animals were observed for dermal irritation approximately 30 minutes after bandage removal and on Days 3, 7, 10, and 14. The

animals were then killed for gross pathological examination.

**Result** There were no deaths. No signs of systemic toxicity or skin irritation

were noted during the study. All animals showed expected gain in bodyweight during the study. No abnormalities were noted at necropsy. The acute dermal median lethal dose (LD50) of the test material in the Sprague-Dawley strain rat was found to be greater than 2000 mg/kg

bodyweight.

**Conclusion** The acute dermal median lethal dose (LD50) of the test material in the

Sprague-Dawley strain rat was found to be greater than 2000 mg/kg

bodyweight.

Reliability (1) valid without restriction

Reference SafePharm Laboratories Limited (1995). Acute Dermal

Toxicity Study in The Rat. Conducted for Chevron Research

and Technology Company, unpublished report.

13.12.2001

**GENETIC TOXICITY 'IN VITRO' (CAS NO.: 151006-60-9)** 

Test substance 1-decene/1-dodecene copolymer, hydrogenated (Alkane 5)

prepared from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer (C50) and higherl and C12 oligomers, CAS 151006-

60-9

Type Salmonella typhimurium and Escherichia coli/Mammalian-Microsome

**Reverse Mutation Assay** 

System of testing Bacterial

**Concentration** 0, 15, 50, 150, 500, 1500, 5000 ug/plate

Cycotoxic conc. Metabolic activation

Result Method

Year GLP > 5000 ug/plate with and without negative

OECD Guide-line 471 "Genetic Toxicology: Salmonella thyphimurium

Reverse Mutation Assay"

1995 Yes

#### **Test condition**

Salmonella typhimurium strains TA1535, TA1537, TA98, and TA100 and Escherichia coli strain WP2uvrA- were treated with the test material using the Ames plate incorporation method at six dose levels, in triplicate, both with and without the addition of a rat liver homogenate metabolizing system (10% liver S9 in standard co-factors). The S9 concentration was determined in an S9 optimization study. The dose range was determined in a preliminary toxicity assay and was 15 to 5000 ug/plate in the first experiment. A second experiment was performed on a separate day using the same dose range as Experiment 1, fresh cultures of the bacterial strains, and fresh chemical formulations. Vehicle (25% w/w Pluronic F127 in ethanol), untreated (negative) and positive controls were included in each experiment.

For the test, 0.1 mL of bacterial culture, 2.0 mL of top agar, 0.1 mL of the test material formulation, vehicle or positive control and either 0.5 mL of S9 mix or phosphate buffer was mixed together and poured onto the surface of a Vogel-Bonner Minimal agar plate. The plates were incubated for 48 hours at 37C after an initial overnight equilibration period and the frequency of revertant colonies was assessed.

For a substance to be considered positive in this test system, it should have induced a dose-related and statistically significant increase in the revertant count (of at least twice the spontaneous reversion rate) in one or more strains of bacteria in the presence and/or absence of S9 in both experiments. To be considered negative, the number of revertants at each dose level should have been less than twofold the vehicle control frequency. Statistical significance was analyzed using the methods recommended by the UKEMS [Reference: Kirkland, D.J., Ed., Statistical Evaluation of Mutagenicity Test Data, UKEMS sub-committee on Guidelines for Mutagenicity Testing. Report Part III (1989) Cambridge University Press.].

### Result

The test material caused no visible reduction in the growth of the bacterial lawn at any dose level either with or without metabolic activation. The test material was therefore tested up to a maximum recommended dose level of 5000 ug/plate. A precipitate was observed at and above 500 ug/plate; this however did not interfere with the scoring of revertant colonies. No significant increase in the frequency of revertant colonies was recorded for any of the bacterial strains with any dose of the test material, either with or without metabolic activation.

The vehicle/suspending agent (Pluronic F127 in ethanol (25% w/w)) and untreated control plates produced counts of revertant colonies within the normal range.

All of the positive control chemicals used in the study induced marked increases in the frequency of revertant colonies, both with and without

the metabolizing system.

The test material was found to be nonmutagenic under the conditions of

this test.

**Conclusion** The test material was found to be nonmutagenic under the conditions of

this test.

**Reliability** (1) valid without restriction

Reference Thompson, P.W. (1995) Salmonella typhimurium and Escherichia

coli/mammalian-microsome reverse mutation assay. Report of Safepharm Laboratories Limited, Derby U.K. conducted for Chevron

Research & Technology Company, Richmond, CA.

31 October 01

## GENETIC TOXICITY 'IN VIVO' (CAS NO.: 151006-60-9)

Test substance 1-decene/1-dodecene copolymer, hydrogenated (Alkane 5)

prepared from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer (C50) and higher] and C12 oligomers. CAS 151006-

60-9

Type Micronucleus assay

Species mouse Sex male/female

Strain CD-1 Route of admin. i.p.

**Exposure period** 24, 48 or 72 hours

**Doses** 1250, 2500 and 5000 mg/kg

**Result** negative

Method OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test"

Year 1995 GLP yes

**Test condition** A study was performed to assess the potential of the test material to

produce damage to chromosomes or aneuploidy when administered via the intraperitoneal route to mice. Following a preliminary range-finding study in males and females, the micronucleus study was conducted using the test material at the maximum recommended dose level of 5000 mg/kg with 2500 and 1250 mg/kg as the lower two dose levels.

In the micronucleus study, groups of ten mice, five males and five females, were given single intraperitoneal doses of the test material at 1250, 2500, and 5000 mg/kg diluted with arachis oil. Further, 4 groups of 10 mice (5 males and 5 females) were dosed, 3 via the intraperitoneal route with arachis oil and one orally with cyclophosphamide to serve as

vehicle and positive controls respectively.

Animals in the vehicle control groups were killed 24, 48 and 72 hours following dosing and positive control group animals were killed 24 hours after exposure. The bone marrow was extracted, and smear preparations were made and stained. The incidence of micronucleated cells per 1000 polychromatic erythrocytes per animal was scored. In addition, the number of normochromatic erythrocytes associated with 1000 erythrocytes were counted; these cells were also scored for incidence of micronuclei.

A positive mutagenic response was demonstrated when a statistically significant and dose responsive increase in the number of micronucleated polychromatic erythrocytes was observed for either the 24, 48, or 72-hour kill times when compared to their corresponding control group. A positive response for bone marrow toxicity was demonstrated when the dose group mean polychromatic to normochromatic ratio was shown to be statistically significantly lower than the concurrent vehicle control group. All data were statistically analysed using appropriate statistical methods as recommended by the UKEMS Sub-committee on Guidelines for Mutagenicity Testing Report, Part III (1989).

Result

There were no premature deaths or clinical signs observed in any of the dose groups. There was no evidence of a statistically significant increase in the frequency of micronucleated polychromatic erythrocytes when compared to the concurrent vehicle control group.

There was no statistically significant change in the PCE/NCE ratio in any of the test material dose groups when compared to their concurrent control groups.

The positive control material produced a marked increase in the frequency of micronucleated polychromatic erythrocytes.

The test material, Alkane 5, was found not to produce an increase in the frequency of micronuclei in polychromatic erythrocytes of mice under the conditions of the test.

Conclusion

The test material, Alkane 5, was considered to be non-genotoxic under the conditions of the test.

Reliability

(1) valid without restriction

Reference

1-decene/1-dodecene copolymer, hydrogenated (Alkane 5) prepared from 10% 1-dodecene and 90% 1-decene, comprising C10 oligomers [approximately 33% trimer (C30), 51% tetramer (C40), 16% pentamer (C50) and higher] and C12 oligomers.

31 October 01

WATER SOLUBILITY (CAS NO.: 163149-28-8)
Test Substance: CAS No.: 163149-28-8;

1-Octene, 1-Decene, 1-Dodecene copolymer, hydrogenated

Method/Guideline: OECD 106

Year (guideline): 1993

Type (test type): Soil Adsorption/Desorption Study

GLP: Yes
Year (study performed): 1995
Temperature: 24 Deg C
pH value: No data
Test Conditions: A saturate

• Note: Concentration prep., vessel type, replication, test

conditions.

A saturated aqueous solution of the test material was prepared by placing 10 mL of ultrapure water (reverse osmosis/ion exchange system) and 4 mL of the test material in a glass tube and rotary-mixing for 16 hours at 24 Deg. C. After settling, the lower aqueous phase was carefully withdrawn using a 10 mL glass syringe.

The water samples were extracted and the extracts were spotted on a Thin Layer Chromatography (TLC) plate, along with a

standard corresponding to 100% recovery of the test material from

water.

Samples were analyzed in duplicate. The method detection limit

was 0.4 mg/L.

**Results:** Water solubility = <0.4 mg/L (less than limit of quantification with

this method).

Units/Value: TLC spots of the duplicate samples were slightly darker than b

 Note: Deviations from protocol or guideline, analytical method. TLC spots of the duplicate samples were slightly darker than blank samples, indicating that a "trace" (<0.4 mg/L) amount of the test material was detected in the saturated water samples, but the concentration was too low to be accurately quantified.

Water solubility was measured as a requirement for a Soil

Adsorption/Desorption (Koc) study, not as part of a water solubility

study.

**Conclusion:** 

**Reliability:** (1) Reliable without restriction

Reference: Stonybrook Laboratories, Inc. 1995. Solubility in Water. Study No.

66471.

Other (source): ExxonMobil Biomedical Sciences, Inc.

## FISH ACUTE TOXICITY (CAS NO.: 163149-28-8)

Test Substance:	CAS No.: 163149-28-8;	
	Octene, 1-Decene, 1-Dodecene copolymer, hydrogenated	
Method/Guideline:	Fish Acute Toxicity Test (EPA 560/6-82-002; OECD 203)	
Type (test type):	Fish Acute Toxicity Test; Dispersion Test	
GLP:	Yes	
Year (study performed):	1994	
Species:	Rainbow Trout (Oncorhynchus mykiss)	

Analytical Monitoring:	Yes (Gas Chromatography - with Flame Ionization Detection)	
Exposure Period:	96 hour	
Statistical Method:	Binomial Probability Analysis (Stephan, et al., 1978)	
Test Conditions:  Note: Concentration prep. vessel type, volume, replication, water quality parameters, environmental conditions, organisms supplier, age, size, weight, loading.	Individual Oil/Water Dispersion (OWD) systems were prepared for each treatment level. The test system was designed to maintain the test substance as a dispersion of small droplets throughout the water column. The test substance was added volumetrically, via graduated cylinder, directly to the dispersion system. Each test chamber was a 10-gallon glass aquaria with 30 liters of water, and was equipped with a vertically mounted, motor-driven impeller assembly. The impeller assembly, consisting of 3-blades on a 10-inch stainless steel shaft, was housed in a 2-inch diameter PVC cylinder with 4 horizontal apertures near the bottom. Water and test substance spilling into the top of the cylinder were expelled through the apertures at the bottom. The OWD systems operated continuously for the duration of the test. Twenty fish were randomly assigned to each chamber. No renewal of the test solutions was performed during the test.  Samples for chemical analysis were removed from the 99 mg/L, 1020 mg/L, and 5010 mg/L concentrations at 10 minutes and 96 hours after test initiation.  Test temperature was 11.7 - 11.8 Deg C., lighting was 16 hours light: 8 hours dark. Dissolved oxygen ranged from 9.6 to 10.1 mg/L and pH ranged from 8.2 to 8.6 during the study. Fish were not fed during the study.  Fish Mean Wt.(Control) = 0.38 g; mean standard length = 3.1 cm; test loading = 0.25 g of fish/L.	
Results:		
Units/Value:	96-hr LL0 = 5010 mg/L, based on nominal loading levels.	
Note: Deviations from protocol or guideline,	Loading Level. Mean Measured Conc. % Mortality @ 96 hr.	
protocol or guideline, analytical method, biological observations, control survival.	Control <loq* 0<="" td=""></loq*>	
	99 mg/L <loq 0<="" td=""></loq>	
	510 mg/L 0	
	1020 mg/L <loq 0<="" th=""></loq>	
	2010 mg/L 0	
	5010 mg/L <loq 0<="" th=""></loq>	
	*LOQ = Limit Of Quantitation = 87 mg/L	

Conclusion:	
Reliability:	Code 1, Reliable without restriction
Reference:	Stonybrook Laboratories, Inc., 1994. Static 96-hour Acute Toxicity Study of MCP-1602 to Rainbow Trout, Study No. 66134.
Other (source):	ExxonMobil Biomedical Sciences, Inc.

# **INVERTEBRATE ACUTE TOXICITY (CAS NO.: 163149-28-8)**

Test Substance:	CAS No.: 163149-28-8;	
	1-Octene, 1-Decene, 1-Dodecene copolymer, hydrogenated	
Method/Guideline:	Daphnia Acute Toxicity Test (EPA 560/6-82-002; OECD 202)	
Type (test type):	Daphnia Acute Toxicity Test	
GLP:	Yes	
Year (study performed):	1994	
Species:	Water flea (Daphnia magna)	
Analytical Monitoring:	Yes (Gas Chromatography - with Flame Ionization Detection)	
Exposure Period:	48-hours	
Statistical Method:	Binomial Probability Analysis (Stephan, et al., 1978)	
Note: Concentration prep.     vessel type, volume,     replication, water quality     parameters, environmental     conditions, organisms     supplier, age, size, loading.	Individual test treatment solutions were prepared as Water Accommodated Fractions (WAFs). A measured amount of the test material was added to 1.0L of test water in aspirator bottles. WAFs were stirred for approximately 20 hours with stir bars producing a vortex of less than 25% of the depth of the solution, and then allowed to settle for approximately 4 hours. Three samples were prepared from each aspirator bottle. For each nominal loading, organisms were added to two labeled 250 ml glass chambers containing 200 ml WAF solution; the third bottle was used for measuring initial water quality. 10 daphnids, less than 24-hours old, were randomly assigned to each test chamber. Test chambers were covered with plexiglass sheets. No renewal of the test solutions was performed during the test.  Test exposure chambers were maintained at 20± 1 °C. Lighting was 16 hours light: 8 hours dark. Dissolved oxygen levels remained above 60% saturation throughout the test. The pH values ranged between 8.0 to 8.08.  Organisms were obtained from in-house cultures.	

Results:	The 48-hr Fl a = 5	5 220 mg/l_hased.or	n nominal loading levels.
Units/Value:	1110 10 111 220 - 0	7,220 mg/2, 5d00d 0.	Themmal loading levels.
Note: Deviations from protocol or guideline, analytical method, biological observations, control survival.			
	Loading Level.	Measured Conc.	% Mortality @ 48 hr.
Results: cont'd	Control	< LOQ*	0
	360 mg/L	< LOQ	0
	630 mg/L	nm	0
	1,350 mg/L	< LOQ	0
	2610 mg/L	nm	0
	5,220 mg/L WAF	< LOQ	0
	*LOQ = Limit Of ( nm = not measure	Quantitation = 2 mg/L ed	-
Conclusion:			
Reliability:	Code 1, Reliable without restriction		
Reference:	Stonybrook Laboratories, Inc. 1994. 48-hour Static Acute Toxicity Study of the WAF of MCP-1602 to <i>Daphnia magna</i> , Study No. 66135.		
Other (source):	ExxonMobil Biomedical Sciences, Inc.		

Algal Toxicity (CAS No.: 163149-28-8)

Test Substance:	CAS No.: 163149-28-8;	
	1-Octene, 1-Decene, 1-Dodecene homopolymer, hydrogenated	
Method/Guideline:	Acute Alga InhibitionTest (EPA 560/6-82-002; OECD 201)	
Type (test type):	Static Acute Alga Inhibition Test	
GLP:	Yes	

Year (study performed):	1994	
Species/Strain:	Green Alga (Selenastrum capricornutum)	
Analytical Monitoring:	Yes (Gas Chromatography - with Flame Ionization Detection)	
Exposure Period:	72 hour	
Statistical Method:	Binomial Probability Analysis (Stephan, et al., 1978)	
Note: Concentration prep. vessel type, volume, replication, water quality parameters, environmental conditions, organism culture, age.	Individual test treatment solutions were prepared as Water Accommodated Fractions (WAFs). Test material was added to 1.0L of algal media in aspirator bottles. The mixing vessels were capped with parafilm and mixed on magnetic stir plates for 20 hours with stir bars producing a vortex of less than 25% of the depth of the solution. After mixing, the solutions were allowed to settle for approximately four hours. The WAF was removed from the bottom of the mixing vessel via the port and used for testing. Test vessels were 125ml glass Erlenmeyer flasks containing 50 mL of treatment solution and algae. Three replicates were prepared for each treatment level. The initial algal cell loading was 1.0 x 10 <sup>4</sup> cells/ml. All test replicates were placed on a shaker table at 100 oscillations per minute during the study. The flasks were covered loosely to facilitate gas exchange. Samples were taken at the end of the 72-hour exposure for cell counts.  Nominal treatment levels were 360, 630, 1350, 2610, and 5220 mg/L. Samples for chemical analysis of the control, 360, 1350, and 5220 mg/L treatment levels were below the Limit of Quantitation (LOQ = 2 mg/L).  Test temperature was 24 ± 1 Deg. C; lighting was continuous at 450 ± 50 foot-candles. The pH was 7.5 at test initiation.	
Results:	70   5  5000 #   1	
Units/Value:	72-hr $EL_0 = 5220$ mg/L, based on biomass.	
Measurement (cells/growth)     Note: Deviations from protocol or guideline, analytical method, biological observations, control survival.	72-hr NOEL = 5220 mg/L, based on biomass.	
Results: cont'd	Mean Cell Loading 72-hr Conc 72 hr Level (mg/L) (% Inhibition) (cells/ml) Control n/a 3.52 x10 <sup>4</sup> 360 12.3 3.09 x10 <sup>4</sup> 630 -12.6* 3.96 x10 <sup>4</sup> 1350 -6.7* 3.75 x10 <sup>4</sup> 2610 -64.0* 5.77 x10 <sup>4</sup> 5220 -76.5* 6.21 x10 <sup>4</sup>	

	n/a - Not applicable *Stimulatory response
Conclusion:	
Reliability:	Code 2, Reliable with Restrictions  Cell growth in the Control did not meet the acceptability requirement of a 16-fold increase. However, growth in the test concentrations indicates that the test material is not inhibitory.
Reference:	Stonybrook Laboratories, Inc. 1994. Static 72-hour Inhibition Study of the WAF of MCP-1602 to Selenastrum capricornutum, Study No. 66137.
Other (source):	ExxonMobil Biomedical Sciences, Inc.

Acute Oral Toxicity (CAS No.: 163149-28-8)

Test Substance	Octene, Decene, Dodecene Copolymer
CAS No.	163149-28-8
Method/Guideline	Other
Type of Study	Acute Oral
GLP	Yes
Year	1995
Species/strain	Sprague-Dawley rats
Sex	Males/Females
No. of animals/sex/dose	5/sex
Route of administration	Oral gavage
Dose/Concentration Levels	Limit dose: 2 g/kg
Remarks on Test Conditions	Five male and five female young adult rats were fasted overnight and dosed via oral gavage with 2 gm/kg of test material. Body weights were recorded prior to fasting and on days 0, 7, and 14. Food was returned to each animal immediately after dosing. Signs of toxicity were recorded approximately 1/2, 1, and 4 hours after dosing and daily thereafter.
D #	
Results	$LD_{50} > 2 g/kg$
Remarks	All animals survived until study termination. Soft stool was noted in two animals at 4 hours and in two animals on day 1.  Chromorhinorrhea was observed in one animal on day 14. No other abnormal clinical signs were observed. There were no gross pathological changes noted at necropsy that were attributed to the test substance.
Conclusions	Under the conditions of this study, octene, decene, dodecene copolymer has a low order of acute toxicity via the oral route of exposure.
Reliability	1 - Reliable without restrictions

Reference	Acute Oral Toxicity in the Sprague-Dawley Rat (1995), Performed by
	Stonybrook Laboratories, Inc.

Repeated Dose Toxicity (CAS No.: 163149-28-8)

Test Substance Octene, Decene, Dodecene Copolymer

CAS No. **163149-28-8** 

Method/Guideline Other

Type of Study Subchronic Dermal

GLP Yes Year 1995

Species/strain Sprague-Dawley rats
Sex Males/Females

No. of animals/sex/dose Route of administration 10/sex/dose Dermal

Dose/Concentration Levels 0, 125, 500, 2000 mg/kg/day

Remarks on Test Conditions | Rats (10/sex/dose) were exposed dermally five days per week for four

weeks at doses of 0, 125, 500, and 2000 mg/kg/day. In addition, two satellite groups (0, 2000 mg/kg/day) were observed for two weeks following the four weeks of dosing to evaluate persistence, delayed effects, and/or recovery. Animals were fitted with Elizabethan collars to minimize ingestion of the test material. The test material was applied to

the skin and left uncovered.

Results NOEL = 500 mg/kg/day

NOAEL = 2000 mg/kg/day

Remarks Treated animals exhibited minimal signs of systemic toxicity. No dermal

irritation was observed at the site of exposure. After the fourth week of dosing, there was a slight decrease in body weight in the males of the high dose group. However, this decrease was statistically significant only for the satellite group of animals. Furthermore, female weight gain was not affected. There were no effects on food consumption during the study. During the last week, male rats of the high dose group had statistically higher segmented neutrophil counts than the control. In addition, males of the high-dose satellite group had significant changes in 7 of 20 serum chemistry parameters. Following the two-week recovery period, no hematologic changes were observed in control or treated animals. After the 2-week recovery period, there were also statistical differences between the female controls and the treated satellite group for 2 serum chemistry parameters. Since these changes in serum chemistry parameters were not consistent between the satellite groups and the matching dose group from the main study, the effects were not considered to be biologically significant. No significant macroscopic changes were noted at the main necropsy or the recovery group necropsy. In addition, no organ weight differences attributable to the test material were noted between the control and treatment groups at either interval. Microscopic changes related to treatment were limited to the skin which showed an increased incidence of hyperplasia and

hyperkeratosis. However, similar changes were also observed in the

32

	control and recovery groups, and thus may be due to repeated shaving of the skin.
Conclusions	Octene, Decene, Dodecene Copolymer has a low order of subchronic toxicity by the dermal route of exposure.
Reliability	2 - Reliable with restrictions – not a guideline study.
Reference	Four-week systemic toxicity study following daily dermal administration to rats, (1995), Mobil Chemical Company.

# Genetic Toxicity 'In Vitro' (CAS No.: 163149-28-8)

Test Substance	Octene, Decene, Dodecene Homopolymer
CAS No.	163149-28-8
Method	Other
Type of Study	Ames
GLP	Yes
Year	1995
Species/Strain	Salmonella typhimurium: TA98, TA100, TA1535, TA1537, TA1538
Metabolic Activation	With and without metabolic activation
Concentrations	0.1, 0.3, 1.0, 3.0, 10.0 μl/50 μl vehicle
Vehicle	Tetrahydrofuran (THF)
Controls	Solvent control and untreated control
Remarks on Test	Serial dilutions of the test substance were prepared in THF to deliver
Conditions	0.1, 0.3, 1.0, 3.0, 10.0 $\mu$ l in 50 $\mu$ l aliquots per bacterial plate. Positive
	controls were 2-aminoanthracene, 9-aminoacridine, 2-nitrofluorene, and
	N-methyl-N'-nitro-N-nitrosoguanidine in DMSO. The S-9 fraction was
	prepared from the livers of Sprague-Dawley rats induced with Aroclor
	1254.
Results	Negative
Remarks for Results	The test meetanial did not induce any taxia offects in the heatania. Name
Remarks for Results	The test material did not induce any toxic effects in the bacteria. None of the strains exhibited reversion frequencies that were substantially
	different (i.e. doubling of frequency or greater) from spontaneous or
	solvent controls in two independent assays. The average spontaneous
	reversion rates for all five strains were also within normal ranges. An
	increased frequency of revertants was observed with the positive
	controls when compared to solvent controls.
	Controls when compared to solvent controls.
Conclusion	MCP-1602 is not mutagenic with or without metabolic activation in this
Conclusion	test system.
	ion of otomic
Reliability	1 - Reliable without restrictions
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Reference	An Ames Salmonella/Mammalian Microsome Mutagenesis Assay (1995)
	Performed by Stonybrook Laboratories, Inc.
Date last changed	September, 01

# **GENETIC TOXICITY 'IN VITRO' (CAS NO.: 163149-28-8)**

Test Substance	Octene, Decene, Dodecene Copolymer
CAS No.	163149-28-8
Method	Other
Type of Study	Chromosomal Aberrations
GLP	Yes
Year	1995
Species/Strain	Cultured Chinese Hamster Ovary (CHO) cells
Metabolic Activation	With and without S9
Concentrations	0.10, 0.20, and 0.40 μl/ml vehicle
Vehicle	Tetrahydrofuran (THF)
Controls	Positive controls: Mitomycin C, cyclophosphamide
Remarks on Test Conditions	Initially, the test material was tested at a range of concentrations (0.0032 $\mu l/ml$ to 0.40 $\mu l/ml$ ) to identify a cytotoxic concentration for the main study. The preliminary assay indicated that the high dose (0.40 $\mu l/ml$ ), which is at or above the limit of solubility of the test substance in the medium, was not toxic. The main study was conducted at the following doses: 0.10, 0.20, and 0.40 $\mu l/ml$ . Cells were exposed to the test chemical for 2 hours in the presence of S9 mix and the cells were harvested 16 hours after initiating treatment. Cells that were not exposed to the S9 fraction were continually exposed to the test substance until cell harvest. The experiment was repeated to confirm the negative findings with the exception that a delayed harvest (40 hours after initiating treatment versus 16 hours) was included.
Results	Negative
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Remarks for Results	Both in the presence an absence of metabolic activation, no significant increase in the proportion of cells with chromosomal aberrations was observed when compared to THF controls. No significant increase in the proportion of cells with chromosomal aberrations compared to the solvent control cultures occurred for either harvest times in assays with or without the S9 fraction. Positive and negative controls in both assays responded as expected.
Conclusion	The test substance is not clastogenic under the conditions of this study.
Reliability	1 - Reliable without restrictions
Reference	Assay for Induction of Chromosomal Aberrations in Cultured Chinese Hamster Ovary (CHO) Cells, (1995). Performed by Stonybrook Laboratories, Inc.
Date last changed	September, 01

Partition Coefficient (CAS No.: 1006-62-1)
Test substance 1-dodecene trimer, hydrogenated (Alkane 4) Test substance

Log pow Method > 7.64 at 20° C

2000

ves

Year GLP

**Test condition** 

The octanol/water partition coefficient was determined using reversephase high performance liquid chromatography (HPLC) with ultraviolet (UV) and refractive index (RI) detectors.

A Bakerbond C-18 column (0.46 cm id x 25 cm, 5 u packing) was used for separation. A mobile phase of 100% acetonitrile (ACN) that allowed UV detection down to 195 nm was used initially. However, a stronger mobile phase of 25% dichloromethane (DCM) and 75% ACN was required. The isocratic (constant mobile phase composition) mode also allowed for sequential use of the RI, a universal detector detector in HPLC. The retention of polycyclic aromatic hydrocarbons (PAHs) standards with known log Kow values was compared to the retention of the PAO to ascertain the log Kow of the PAO.

Result

The retention times of the polyalphaolefin (PAO), Alkane 4, and nine polycyclic aromatic hydrocarbons (PAHs) with known log Kow values were compared. Neither the ultraviolet (UV) nor refractive index (RI) detectors saw any evidence of PAO elution. The standard mixture of PAHs that eluted had log Kow values of up to 7.64. Therefore, the PAO log Kow value is greater than 7.64 but can be extrapolated to greater than 8. Since coronene eluted at 14.2 minutes and no PAO was detected during the 70 minute analysis time, it can be reasonably concluded that the log Kow of PAO is greater than 8. This high value is emphasized by its limited solubility in the polar solvent acetonitrile. The required use of MTBE implies a very high organic phase preference.

Conclusion

The PAO, Alkane 4, log Kow value is greater than 7.64 but can be extrapolated to greater than 8.

(1) valid without restriction

Reliability Reference

Seary, M. (2000) Determination of Water Solubility and Octanol/Water Partition Coefficient of C-12 Trimer Polyalphaolefin. Unpublished report conducted by Integrated Laboratory Technologies, Chevron Research and Technology Co., Richmond, CA.

14.03.2001

Water Solubility (CAS 151006-62-1)

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Value < 1 other: ppt at  $^{\circ}$  C Qualitative insoluble (< 0.1 mg/L)

Pka at 25 ° C Ph at and ° C

Method

Year 2000

**GLP** Yes

**Test condition** A low-pressure chromatographic technique was used for concentrating the

polyalphaolefin (PAO) to determine its solubility in water. A generator column was prepared with an inert packing material coated with PAO. High-purity water was slowly percolated through the column. The effluent, containing any dissolved PAO, then passed through an octadecyl-bonded phase (C-18) silica collection column. This column retained any organic material from the polar aqueous medium. Acetonitrile and dichloromethane washes were used to elute any PAO from the collector column. The remaining residue was analyzed by gas chromatography-mass spectrometry (GC-MS) to determine the amount of PAO in this fraction.

The water solubility of PAO was calculated based on the total volume of

water passed through the columns.

**Result** The results of the test showed that no polyalphaolefin (PAO) was

observed. The only material seen was C-18 carboxylic acid arising from degradation of some of the bonded phase on the collection column. The limit of detection for PAO, <10 ng, was calculated from comparison to injections of standard solutions, pyrene, perylene and coronene.

Therefore, the water solubility was calculated to be <1 ppt.

**Conclusion** The water solubility of the polyalphaolefin, Alkane 4, was calculated to be

<1 ppt.

**Reliability** (1) valid without restriction

**Reference** Seary, M. (2000) Determination of Water Solubility and Octanol/Water

Partition Coefficient of C-12 Trimer Polyalphaolefin. Unpublished report conducted by Integrated Laboratory Technologies, Chevron Research and

Technology Co., Richmond, CA.

14.03.2001

### ACUTE/PROLONGED TOXICITY TO FISH (CAS NO.: 151006-62-1)

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

**Type** semistatic

**Species** Oncorhynchus mykiss (Fish, fresh water)

 Exposure period
 96 hour(s)

 Unit
 mg/l

 Analytical monitoring
 yes

 NOEC
 >= 1000

 LC50
 > 1000

Method OECD Guide-line 203 "Fish, Acute Toxicity Test"

Year 1995 GLP Yes

## **Test condition**

A study was performed to assess the acute toxicity of the test material, Alkane 4, to rainbow trout. Following a preliminary range-finding study, fish were exposed, in two groups of ten, to a Water Accommodated Fraction (WAF) of the test material for a period of 96 hours. A semi-static test regime was employed in the study involving a daily renewal of the test preparations to ensure that the concentrations of the test material remained near nominal and to prevent the build up of nitrogenous waste products. The WAF was prepared by placing the test material on the surface of water to give a 1000 mg/L loading rate which was then stirred with a magnetic stirrer to achieve a vortex depth of approximately 20-25% of the distance to the bottom of the vessel for 24 hours. The mixture was then allowed to stand for 4 hours prior to removing the aqueous phase or WAF by siphon.

The number of mortalities and any adverse reactions to exposure in each test and control vessel were determined 3 and 6 hours after the start of exposure and then daily throughout the study until termination after 96 hours. Duplicate control groups were maintained under identical conditions but not exposed to the test material. The vessels received no auxiliary aeration and were covered to reduce evaporation.

#### Result

In the Range-finding study the results showed no mortalities at the 100 and 1000 mg/L loading rate Water Accommodated Fractions (WAF's).

The results of the definitive study showed the highest loading rate WAF resulting in 0% mortality to be greater than or equal to 1000 mg/L, the lowest loading rate WAF resulting in 100% mortality to be greater than 1000 mg/L and the No Observed Effect Concentration (NOEC) to be greater than or equal to 1000 mg/L loading rate WAF. The No Observed Effect Concentration is based upon zero mortalities and the absence of any adverse effects of exposure at this concentration.

Analysis of the WAF was carried out by Total Organic Carbon (TOC) analysis. Water samples were taken from the control and each replicate test vessel at 0 hours (fresh test preparations), 24 hours (old test preparations), 72 hours (fresh test preparations) and 96 hours (old test preparations). The results of the TOC analysis showed that, compared to the controls, no significant levels of carbon were detected in the WAFs.

## Conclusion

The 96-hour median Lethal Loading Rate (LLR50) for the test material to rainbow trout (Oncorhynchus mykiss), based on nominal loading rates, was greater than 1000 mg/L loading rate Water Accommodated Fraction and correspondingly the No Observed Effect Concentration was greater than or equal to 1000 mg/L loading rate Water Accommodated Fraction.

# Reliability

(1) valid without restriction

## Reference

SafePharm Laboratories Limited (1995). Acute Toxicity to Rainbow Trout. Conducted for Chevron Research and Technology Company, unpublished report.

27.02.2001

# **ACUTE TOXICITY TO AQUATIC INVERTEBRATES (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type static

Species Daphnia magna (Crustacea)

Exposure period 48 hour(s)
Unit mg/l
Analytical monitoring yes
NOEC >= 1000
EC50 > 1000

Method OECD Guide-line 202, part 1 "Daphnia sp., Acute Immobilisation Test"

Year 1995 GLP Yes

## **Test condition**

A study was performed to assess the acute toxicity of the test material, Alkane 4, to Daphnia magna. Following a preliminary range-finding study, forty daphnids (4 replicates of 10 animals) were exposed to a Water Accommodated Fraction (WAF) of the test material for 48 hours under static test conditions. The WAF was prepared by placing the test material on the surface of the water to give a 1000 mg/L loading rate which was then stirred by magnetic stirrer to achieve a vortex depth of approximately 20-25% of the distance to the bottom of the vessel for 24 hours. The mixture was then allowed to stand for 4 hours prior to removing the aqueous phase or WAF by siphon. Immobilization and any adverse reactions to exposure were recorded after 24 and 48 hours. Replicate control groups were maintained under identical conditions but not exposed to the test material. The vessels received no auxiliary aeration and were covered to reduce evaporation.

#### Result

In the Range-finding study the results showed no immobilization at the 100, and 1000 mg/L loading rate Water Accommodated Fractions (WAF).

In the Definitive study, there was no immobilization in 40 daphnids exposed to a 1000 mg/L loading rate WAF for a period of 48 hours.

The No Observed Effect Concentration after 24 and 48 hours exposure was greater than or equal to 1000 mg/L loading rate WAF. The No Observed Effect Concentration is based upon zero immobilization at this concentration.

Analysis of the Water Accommodated Fractions was carried out by Total Organic Carbon (TOC) analysis on the test preparation at 0 and 48 hours. The results of the TOC analysis showed that compared to the controls, no significant levels of carbon were detected in the WAFs.

#### Conclusion

The No Observed Effect Concentration after 24 and 48 hours exposure was greater than or equal to 1000 mg/L loading rate WAF. The No Observed Effect Concentration is based upon zero immobilization at this concentration.

**Reliability** (1) valid without restriction

**Reference** SafePharm Laboratories Limited (1995). Acute Toxicity to Daphnia Magna. Conducted for Chevron Research and Technology Company,

27.02.2001

## **TOXICITY TO AQUATIC PLANTS E.G. ALGAE (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Species Selenastrum capricornutum (Algae)

Endpoint growth rate
Exposure period 96 hour(s)
Unit mg/l
Analytical monitoring yes
NOEC >= 1000
EC50 > 1000

Method OECD Guide-line 201 "Algae, Growth Inhibition Test"

Year 1995 GLP Yes

#### **Test condition**

A study was performed to assess the effect of the test material, Alkane 4, on the growth of Selenastrum capricornutum. Following a preliminary range-finding study, Selenastrum capricornutum was exposed to a Water Accommodated Fraction (WAF) of the test material (six replicate flasks) for 96 hours under constant illumination and shaking at a temperature of 24oC. The WAF was prepared by placing the test material on the surface of the water to give a 1000 mg/L loading rate which was then stirred to achieve a vortex depth of approximately 20-25% of the distance to the bottom of the vessel for 24 hours. The mixture was then allowed to stand for 4 hours prior to removing the aqueous phase or WAF by siphon. Samples of the algal populations were removed daily, and algal cell concentrations were determined, using an electronic cell counter, for each control and treatment group. Two replicate control groups were maintained under identical conditions but not exposed to the test material.

At the initiation of the study, the algal suspension culture contained a nominal cell density of 10,000 cells per mL.

A Student's t-test was carried out on the area under the growth curve data at 96 hours for the control and 1000 mg/L loading rate WAF test concentration to determine any statistically significant differences between the test and control groups.

#### Result

In the Range-finding study the results showed no effect on growth at 1000 mg/L Water Accommodated Fraction (WAF).

From the results of the definitive study neither the growth or the biomass of Selenastrum capricornutum were affected by the presence of the test material over the 96-hour exposure period.

All test and control cultures were inspected microscopically at 96 hours. There were no abnormalities detected in any of the control or test cultures.

Analysis of the WAF was carried out by Total Organic Carbon (TOC) analysis on samples from two replicate vessels of treated and control media

at the beginning and end of the test. Given the background level of carbon in the control vessels and also the low level of carbon in the test vessels it is considered that all the results were around the limit of detection of the analytical method. Accordingly the results do not provide definite evidence of stability of the test preparations.

The effect of Alkane 4 on the growth of Selenastrum capricornutum has been investigated and gave median effective loading rate (ELR50) values of greater than 1000 mg/L loading rate WAF. Correspondingly the No Observed Effect Concentration was greater than or equal 1000 mg/L loading rate WAF. These results are based on an initial loading rate of 2000 mg/L which was diluted by the addition of the algal suspension to give an equivalent loading rate of 1000 mg/L.

**Conclusion** Exposure of Selenastrum capricornutum to the test material gave median

Effective Loading Rate (ELR50) values of greater than 1000 mg/L loading rate WAF and correspondingly the No Observed Effect Concentration was

greater than or equal 1000 mg/L loading rate WAF.

Reliability (1) valid without restriction

Reference SafePharm Laboratories Limited (1995). Algal Inhibition Test. Conducted for

Chevron Research and Technology Company, unpublished report.

27.02.2001

# **TOXICITY TO MICROORGANISMS E.G. BACTERIA (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type aquatic

**Species** activated sludge of a predominantly domestic sewage

Exposure period 3 hour(s)
Unit mg/l
Analytical monitoring no
EC50 > 1000
NOEC >= 1000

Method OECD Guide-line 209 "Activated Sludge, Respiration Inhibition Test"

Year 2000 GLP Yes

## **Test condition**

A study was performed to assess the effect of the test material on the respiration of activated sewage sludge.

Following a preliminary range-finding study, activated sewage sludge was exposed to an aqueous dispersion of the test material at a concentration of 1000 mg/L (three replicate flasks) for a period of 3 hours at 21oC with the addition of a synthetic sewage as a respiratory substrate. 500 mg of test material was dispersed directly into approximately 250 ml of water. Synthetic sewage (16 ml), activated sewage sludge (200 ml) and water were added to a final volume of 500 ml to give the required concentration of 1000 mg/L.

The rate of respiration was determined after 30 minutes and 3 hours contact time and compared to data for the control and a reference material, 3,5-dichlorophenol.

Observations were made on the test preparations throughout the study period, and the pH of the control, reference material and test material preparations was measured at the end of the exposure period prior to measurement of the oxygen consumption rate.

The results of the study were considered valid if (1) the two control respiration rates are within 15% of each other and (2) the EC50 (3-hour contact time) for 3,5-dichlorophenol lies within the range 5 to 30 mg/L.

#### Result

Alkane 4 EC50 (3hours): >1000 mg/L 3,5-dichlorophenol EC50 (3hours): 11 mg/L Variation in respiration rates of controls 1 and 2 after 3 hours: +/- 3%.

The No Observed Effect Concentration (NOEC) after 3 hours was 1000 mg/L.

The validation criteria for the control respiration rates and reference material EC50 values were satisfied.

It was considered unnecessary and unrealistic to test at concentrations in excess of 1000 mg/L.

#### Conclusion

The effect of the test material on the respiration of activated sewage sludge micro-organisms gave a 3-hour EC50 of greater than 1000 mg/L. The No Observed Effect Concentration (NOEC) after 3 hours exposure was 1000 mg/L.

## Reliability

(1) valid without restriction

#### Reference

SafePharm Laboratories Limited (2000). Assessment of the Inhibitory Effect of the Respiration of Activated Sewage Sludge. Conducted for Chevron Research and Technology Company, unpublished report.

13.03.2001

# **ACUTE ORAL TOXICITY (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type LD50 Species rat

Strain Sprague-Dawley
Sex male/female

Number of animals 10

Vehicle other: none value > 5000 mg/kg bw

Method OECD Guide-line 401 "Acute Oral Toxicity"

Year 1995 GLP Yes

**Test condition** A study was performed to assess the acute oral toxicity of the test material

in the Sprague-Dawley strain rat. Following a range-finding study, a group of ten fasted animals (five males and five females) was given a single oral dose of undiluted test material at a dose level of 5000 mg/kg bodyweight. Individual bodyweights were recorded on the day of dosing to allow calculation of individual treatment volumes and on Days 7 and 14. The animals were observed for deaths or overt signs of toxicity 1, 2.5, and 4 hours after dosing and subsequently once daily for 14 days. They were

then killed and subjected to a gross necropsy.

**Result** There were no deaths. No signs of systemic toxicity were noted during the

study. All animals showed expected gain in body weight during the study. No abnormalities were noted at necropsy. The acute oral median lethal dose (LD50) of the test material in the Sprague-Dawley strain rat was

found to be greater than 5000 mg/kg bodyweight.

**Conclusion** The acute oral median lethal dose (LD50) of the test material in the

Sprague-Dawley strain rat was found to be greater than 5000 mg/kg

bodyweight.

**Reliability** (2) valid without restriction

Reference SafePharm Laboratories Limited (1995). Acute Oral Toxicity Study in The

Rat. Conducted for Chevron Research and Technology Company,

unpublished report.

12.03.2001

# **ACUTE INHALATION TOXICITY (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type LC50 Species rat

Strain Sprague-Dawley
Sex male/female

Number of animals 10

Vehicleother: noneExposure time4 hour(s)Value> 5.06 mg/l

Method OECD Guide-line 403 "Acute Inhalation Toxicity"

Year 1995 GLP Yes

#### **Test Condition**

A study was performed to assess the acute inhalation toxicity of the test material, as supplied, by exposing a single group of Sprague-Dawley strain rats (five males and five females) to an aerosol atmosphere. The animals were exposed for four hours using a nose only exposure system.

Prior to the start of the study, test material atmospheres were generated within the exposure chamber. During these periods air flow settings, test material input and the sampling system were varied to achieve the required atmospheric concentrations. During the exposure period, temperature, relative humidity, oxygen concentrations and nominal atmospheric concentrations were monitored at regular intervals. The particle size of the generated atmosphere of the test material inside the exposure chamber was determined four times during the exposure period using a Cascade Impactor.

Clinical observations were performed hourly during the exposure, immediately at the end of the exposure, one hour after the termination of the exposure and once daily for 14 days. Individual bodyweights were recorded on the day of exposure and on Days 7 and 14. Necropsies were performed on all animals at study termination.

## Result

The mean achieved atmosphere concentration was 5.06 mg/L. The mean mass median aerodynamic diameter was 1.2 u. The inspirable fraction (%<4 u) was 90.1%. The geometric standard deviation was 0.40u. No deaths occurred. Common abnormalities noted during the study were wet fur, hunched posture, and piloerection. Incidents of increased respiratory rate and ptosis and isolated incidents of decreased respiratory rate and red/brown staining on the head were noted. All animals appeared normal on Day 2 following exposure and for the remainder of the study. Normal bodyweight gain was noted during the study. No abnormalities were detected at necropsy. The acute inhalation median lethal concentration (LC50) of the test material, Alkane 4, in the Sprague-Dawley strain rat was greater than 5.06 mg/L.

## Conclusion

The acute inhalation median lethal concentration (LC50) of the test material, Alkane 4, in the Sprague-Dawley strain rat was greater than 5.06 mg/L.

## Reliability

(2) valid without restriction

# Reference

SafePharm Laboratories Limited (1995). Acute Inhalation Toxicity Study (Nose Only) in The Rat. Conducted for Chevron Research and Technology Company, unpublished report.

12.03.2001

# **ACUTE DERMAL TOXICITY (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type LD50 Species rat

Strain Sprague-Dawley
Sex male/female

Number of animals 10

Vehicle other: none Value > 2000 mg/kg bw

Method OECD Guide-line 402 "Acute dermal Toxicity"

Year 1995 GLP Yes

Test condition

A study was performed to assess the acute dermal toxicity of the test material in the Sprague-Dawley strain rat. A group of ten animals (five males and five females) was given a single, 24-hour, semi-occluded, dermal application to intact skin at a dose level of 2000 mg/kg bodyweight. Individual bodyweights were recorded on the day of dosing to allow calculation of individual treatment volumes and on Days 7 and 14. The animals were observed for deaths or overt signs of toxicity 1, 2.5, and 4 hours after dosing and subsequently once daily for 14 days. The animals were observed for dermal irritation approximately 30 minutes after bandage removal and on Days 3, 7, 10, and 14. The

animals were then killed for gross pathological examination.

**Result** There were no deaths. No signs of systemic toxicity or skin irritation

were noted during the study. All animals showed expected gain in bodyweight during the study. No abnormalities were noted at necropsy. The acute dermal median lethal dose (LD50) of the test material in the Sprague-Dawley strain rat was found to be greater than 2000 mg/kg

bodyweight.

**Conclusion** The acute dermal median lethal dose (LD50) of the test material in the

Sprague-Dawley strain rat was found to be greater than 2000 mg/kg

bodyweight.

**Reliability** (2) valid without restriction

Reference SafePharm Laboratories Limited (1995). Acute Dermal Toxicity Study in

The Rat. Conducted for Chevron Research and Technology Company,

unpublished report.

22.02.2001

# REPEATED DOSE TOXICITY (CAS NO.: 151006-62-1)

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

**Species** rat

Sexmale/femaleStrainSprague-Dawley

Route of admin. unspecified 28 days Frequency of treatment Post obs. period 14 days Doses 0, 1000

**Control group** yes, concurrent no treatment

**NOAEL** = 1000 mg/kg bw

Method OECD Guide-line 407 "Repeated Dose Oral Toxicity - Rodent: 28-day

or 14-d Study"

Year 1995 GLP Yes

## **Test condition**

The test material was administered by gavage to a group of five male and five female Sprague-Dawley CD strain rats for twenty-eight consecutive days at a dose level of 1000 mg/kg/day. A control group of five males and five females remained untreated throughout the study period but was otherwise handled in an identical manner to the test animals. Two satellite groups, each of five males and five females, were similarly treated at 1000 mg/kg/day or remained untreated respectively; satellite groups were maintained without treatment for a further fourteen days following the end of the dosing period. Clinical signs, bodyweight, and food and water consumption were monitored during the study. Hematology and blood chemistry were evaluated for all main group animals during the final week of dosing and for satellite group animals at the end of the treatment-free period. All animals were subjected to a gross necropsy examination. The following organs were weighed: adrenals, brain, gonads, heart, kidneys, liver, pituitary, spleen. Histopathological evaluation of the following tissues from main test and control animals was performed: adrenals, heart, kidneys, liver, spleen, testes, macroscopically observed lesions.

## Result

There were no deaths during the study. No clinically observable signs of toxicity were detected in test or control animals throughout the study period. No adverse effect on bodyweight development was detected. No adverse effect on dietary intake was detected. No overt intergroup differences in water consumption were detected. No treatment-related effects were detected for hematology, blood chemistry, or organ weights. No treatment-related macroscopic abnormalities were detected at necropsy, and no treatment-related microscopic changes were observed.

Oral administration of the test material, Alkane 4, to rats for a period of twenty-eight consecutive days at a dose level of 1000 mg/kg/day produced no treatment-related changes in the parameters measured. The "No Observed Effect Level" (NOEL) is therefore considered to be 1000 mg/kg/day.

## Conclusion

Oral administration of the test material, Alkane 4, to rats for a period of twenty-eight consecutive days at a dose level of 1000 mg/kg/day produced no treatment-related changes in the parameters measured. The "No Observed Effect Level" (NOEL) is therefore considered to be 1000 mg/kg/day.

## Reliability

(1) valid without restriction

# **Reference** SafePharm Laboratories Limited (1995). Twenty-Eight Day Sub-Acute

Oral (Gavage) Toxicity Study in the Rat - Limit Test, Including Recovery Groups. Conducted for Chevron Research and Technology Company,

unpublished report.

28.02.2001

# **GENETIC TOXICITY 'IN VITRO' (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

**Type** other: Salmonella typhimurium and Escherichia coli/Mammalian-

Microsome Reverse Mutation Assay

System of testing Bacterial

**Concentration** 0, 15, 50, 150, 500, 1500, 5000 ug/plate

Cycotoxic conc. > 5000 ug/plate
Metabolic activation with and without
Result negative

Method OECD Guide-line 471 "Genetic Toxicology: Salmonella thyphimurium

Reverse Mutation Assay"

Year 1995 GLP Yes

#### **Test condition**

Salmonella typhimurium strains TA1535, TA1537, TA98, and TA100 and Escherichia coli strain WP2uvrA- were treated with the test material using the Ames plate incorporation method at six dose levels, in triplicate, both with and without the addition of a rat liver homogenate metabolizing system (10% liver S9 in standard co-factors). The dose range was determined in a preliminary toxicity assay and was 15 to 5000 ug/plate in the first experiment. A second experiment was performed on a separate day using the same dose range as Experiment 1, fresh cultures of the bacterial strains, and fresh chemical formulations. Vehicle (25% w/w Pluronic F127 in ethanol), untreated (negative) and positive controls were included in each experiment.

For the test, 0.1 mL of bacterial culture, 2.0 mL of top agar, 0.1 mL of the test material formulation, vehicle or positive control and either 0.5 mL of S9 mix or phosphate buffer was mixed together and poured onto the surface of a Vogel-Bonner Minimal agar plate. The plates were incubated for 48 hours at 37C after an initial overnight equilibration period and the frequency of revertant colonies was assessed.

For a substance to be considered positive in this test system, it should have induced a dose-related and statistically significant increase in the revertant count (of at least twice the spontaneous reversion rate) in one or more strains of bacteria in the presence and/or absence of S9 in both experiments. To be considered negative, the number of revertants at each dose level should have been less than twofold the vehicle control frequency. Statistical significance was analyzed using the methods recommended by the UKEMS [Reference: Kirkland, D.J., Ed., Statistical Evaluation of Mutagenicity Test Data, UKEMS sub-committee on Guidelines for Mutagenicity Testing. Report Part III (1989) Cambridge University Press.].

The test material formulations were assessed for concentration, stability and homogeneity. The formulations were shown to be stable and the concentrations were found to be within +/-10% of nominal.

#### Result

The test material caused no visible reduction in the growth of the bacterial lawn at any dose level either with or without metabolic activation. The test material was therefore tested up to a maximum recommended dose level of 5000 ug/plate. A precipitate was observed at and above 1500 ug/plate; this however did not interfere with the scoring of revertant colonies. No significant increase in the frequency of revertant colonies was recorded for any of the bacterial strains with any dose of the test material, either with or without metabolic activation.

The vehicle/suspending agent (Pluronic F127 in ethanol (25% w/w)) and untreated control plates produced counts of revertant colonies within the normal range.

All of the positive control chemicals used in the study induced marked increases in the frequency of revertant colonies, both with and without the metabolizing system.

The test material was found to be nonmutagenic under the conditions of this test.

Conclusion

The test material was found to be nonmutagenic under the conditions of

this test.

**Reliability** (1) valid without restriction

**Reference** SafePharm Laboratories Limited (1995). Salmonella typhimurium and

Escherichia coli/Mammalian-Microsome Reverse Mutation Assay. Conducted for Chevron Research and Technology Company,

unpublished report.

12.03.2001

# **GENETIC TOXICITY - IN VITRO (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type Mammalian Cell Gene Mutation Test

System of testing CHO HGPRT

**Concentration** 313, 625, 1250, 2500, 5000 ug/ml

Cycotoxic conc. > 5000 ug/ml Metabolic activation with and without

**Result** negative

Method OECD Guide-line 476 "Genetic Toxicology: In vitro Mammalian Cell

Gene Mutation Test"

Year 2001 GLP Yes

**Test condition** Chinese hamster ovary cells treated with the test material were

evaluated for gene mutations at the HGPRT locus with duplicate

cultures, together with vehicle (ethanol) and positive controls.

A rangefinding test was conducted to assess toxicity of the test article to the cells. In the non-activated system, concentrations of 0.5 to 5000 ug/mL had relative cloning efficiencies (RCE) ranging from 97% to 73%. In the activated system, the RCEs ranged from 122% to 80%.

In Experiment 1, dose levels of 313, 625, 1250, 2500 and 5000 ug/mL were tested. Cultures tested with and without an induced rat liver homogenate metabolizing system (10% S9 in standard co-factors, 2% S9 final concentration) were exposed for 4 hours, after which the cells were washed, trypsinized and seeded for parallel cytotoxicity and mutant expression determination. For cytotoxicity, cells were seeded in 3 plates for each replicate at a density of 200 cells/60 mm dish and incubated for 7 days. For expression of 6-thioguanine (TG)-resistant mutants (HGPRT locus), cells were subcultured at a density of 2x10<sup>6</sup> cells/150 cm<sup>2</sup> flask and subcultured at 2- to 3-day intervals for a period of 9 days. After the expression period, the cells from each treated replicate were harvested and seeded in 12 100 mm plates at a density of 2x10<sup>5</sup> cells/plate. To determine the cloning efficiency of the cells at the time of selection, 200 cells/60 mm dish were plated in triplicate in cloning medium. All clonable test doses and positive and solvent controls were cloned for mutant selection. Cultures were incubated for 7 days. After the incubation period, the colonies were washed, fixed, stained, and counted for cloning efficiency and mutant selection.

Results were confirmed in a second experiment using the same procedures and dose levels.

Results were analyzed by the Cochran-Armitage test for trend and the Fisher-Irwin exact test for group comparisons for proportions. Within group comparisons were made by the Fisher-Irwin exact test.

The test results were considered to have caused a positive response if the test article yielded an average mutant frequency greater than 15 mutants per 1x10<sup>6</sup> surviving cells and showed more than a significant (statistically or 2-fold) increase in the number of mutants per 1-10<sup>6</sup> surviving cells over that of the concurrent and historical solvent controls. In the absence of a positive dose-response trend, at least 2 consecutive test doses must have shown a significant increase in the number of mutants.

All vehicle (solvent) controls gave frequencies of mutations within the range expected. All the positive control treatments gave significant increases in mutant frequency indicating the satisfactory performance of the test and of the activity of the metabolizing system. In the first mutation assay, the RCEs ranged from 92% to 77% and 111% to 89% with and without metabolic activation, respectively. In the confirmatory assay, RCEs ranged from 50% to 23% and 89% to 52% with and without activation, respectively.

In the first experiment with activation, at 625 ug/mL, a significant increase in mutant frequency was seen compared to the solvent control [4, 4, 9, 4, 5, 2 mutants/10<sup>6</sup> surviving cells at 0, 313, 625, 1250, 2500, and 5000 ug/mL, respectively]; however, the increase was not significant when compared to the historical control [9 mutants/10<sup>6</sup> surviving cells].

Result

The same was true for the 2500 ug/mL dose level with activation in the confirmatory assay: 1, 3, 1, 3, 8, 3 mutants/10<sup>6</sup> surviving cells at 0, 313, 625, 1250, 2500, and 5000 ug/mL, respectivelyl.

The positive controls caused a significant increase in mutant frequencies and all criteria for a valid assay were met.

The test material was considered to be negative in this assay.

**Conclusion** The test material was shown to be non-mutagenic when tested under the

conditions of this assay.

**Reliability** (1) valid without restriction

Reference SITEK Research Laboratories (2001). Test for Chemical Induction of

Gene Mutation at the HGPRT Locus in Cultured Chinese Hamster Ovary (CHO) Cells With and Without Metabolic Activation With a Confirmatory Assay. Conducted for Chevron Research and Technology Company,

unpublished report.

12-10-2001

# **GENETIC TOXICITY - IN VITRO (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

Type Chromosomal aberration test

System of testing Human Lymphocyte

**Concentration** 39, 78.1, 156.25, 312.5, 625, 1250, 2500, 5000 ug/ml

Cycotoxic conc. > 5000 ug/ml Metabolic activation with and without

**Result** negative

Method OECD Guide-line 473 "Genetic Toxicology: In vitro Mammalian

Cytogenetic Test"

Year 1995 GLP Yes

**Test condition** Human lymphocytes treated with the test material were evaluated for

chromosome aberrations with duplicate cultures, together with vehicle

(ethanol) and positive controls.

In Experiment 1, 8 dose levels ranging from 39 to 5000 ug/ml were tested. Cultures with an induced rat liver homogenate metabolizing system (10% S9 in standard co-factors) were exposed for 4 hours, after which the medium was replaced and the cultures were re-incubated for a further 16 hours. Cultures without metabolic activation were treated continuously for 20 hours. Mitotic indices demonstrated that there was no toxicity.

Therefore, the top three dose levels (1250, 2500, and 5000 ug/ml) were

evaluated for chromosome aberrations and polyploidy.

Results were confirmed in a second experiment. The cultures with metaboloc activation were treated for 4 hours and harvested 16 hours (concentrations of 625, 1250, 2500, 5000 ug/ml) and 40 hours (1250, 2500, 5000 ug/ml) later. Positive controls were evaluated only in the 20hour harvest cultures. The dose levels selected for metaphase analysis (1250, 2500, and 5000 ug/ml for the 20hr harvests and 5000 ug/ml for the 44 hr harvests) were selected on the basis of toxicity demonstrated by the mitotic index. Slides were coded and blindly scored. A total of 2000 lymphocyte cell nuclei were counted and the number of cells in metaphase recorded and expressed as the mitotic index and as a percentage of the vehicle control value. Where possible, the first 100 consecutive wellspread metaphases from each culture were counted, and if the cell had 46 or more chromosomes, any gaps, breaks or rearrangements were noted. The frequency of cells with aberrations (both including and excluding gaps) and the frequency of polyploid cells was compared, where necessary, with the concurrent vehicle control value using Fisher's Exact test.

The test material formulations were assessed for concentration, stability and homogeneity. The formulations were shown to be stable and the concentrations were found to be within +/-10% of nominal.

Result

All vehicle (solvent) controls gave frequencies of cells with aberrations within the range expected for normal human lymphocytes. All the positive control treatments gave significant increases in the frequency of cells with aberrations indicating the satisfactory performance of the test and of the activity of the metabolizing system. The test material, induced no statistically significant increases in the frequency of cells with aberrations or polyploid cells. The test material was shown to be nonclastogenic to human lymphocytes in vitro.

Conclusion

The test material was shown to be nonclastogenic to human lymphocytes

in vitro.

Reliability

(2) valid without restriction

Reference

SafePharm Laboratories Limited (1995). Chromosome Aberration Test in Human Lymphocytes. Conducted for Chevron Research and Technology Company, unpublished report.

12.03.2001

# **GENETIC TOXICITY 'IN VIVO' (CAS NO.: 151006-62-1)**

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

**Type** Micronucleus assay

Species mouse
Sex male/female

Strain CD-1 Route of admin. i.p.

**Exposure period** 24, 48 or 72 hours

**Doses** 1250, 2500 and 5000 mg/kg

**Result** negative

Method Year GLP OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test" 1995 yes

#### **Test condition**

A study was performed to assess the potential of the test material to produce damage to chromosomes or aneuploidy when administered via the intraperitoneal route to mice. Following a preliminary range-finding study in males and females, the micronucleus study was conducted using the test material at the maximum recommended dose level of 5000 mg/kg with 2500 and 1250 mg/kg as the lower two dose levels.

In the micronucleus study, groups of ten mice, five males and five females, were given single intraperitoneal doses of the test material at 1250, 2500, and 5000 mg/kg diluted with arachis oil. Further, 4 groups of 10 mice (5 males and 5 females) were dosed, 3 via the intraperitoneal route with arachis oil and one orally with cyclophosphamide to serve as vehicle and positive controls respectively.

Animals in the vehicle control groups were killed 24, 48 and 72 hours following dosing and positive control group animals were killed 24 hours after exposure. The bone marrow was extracted, and smear preparations were made and stained. The incidence of micronucleated cells per 1000 polychromatic erythrocytes per animal was scored. In addition, the number of normochromatic erythrocytes associated with 1000 erythrocytes were counted; these cells were also scored for incidence of micronuclei.

A positive mutagenic response was demonstrated when a statistically significant and dose responsive increase in the number of micronucleated polychromatic erythrocytes was observed for either the 24, 48, or 72-hour kill times when compared to their corresponding control group. A positive response for bone marrow toxicity was demonstrated when the dose group mean polychromatic to normochromatic ratio was shown to be statistically significantly lower than the concurrent vehicle control group. All data were statistically analysed using appropriate statistical methods as recommended by the UKEMS Sub-committee on Guidelines for Mutagenicity Testing Report, Part III (1989).

## Result

There were no premature deaths or clinical signs observed in any of the dose groups. There was a small statistically significant increase in the frequency of micronucleated polychromatic erythrocytes in the 24-hour 5000 mg/kg test material dose group when compared to the concurrent vehicle control group. The response was not part of a dose-related effect, was within the current historical range for 24-hr vehicle control groups, and was, therefore considered to be spurious and of no toxicological significance.

There was no statistically significant change in the PCE/NCE ratio in any of the test material dose groups when compared to their concurrent control groups.

The positive control material produced a marked increase in the frequency of micronucleated polychromatic erythrocytes.

The test material, Alkane 4, was found not to produce a toxicologically significant increase in the frequency of micronuclei in polychromatic erythrocytes of mice under the conditions of the test.

**Conclusion** The test material, Alkane 4, was found not to produce a toxicologically

significant increase in the frequency of micronuclei in polychromatic

erythrocytes of mice under the conditions of the test.

**Reliability** (1) valid without restriction

**Reference** SafePharm Laboratories Limited (1995). Micronucleus Test in the Mouse.

Conducted for Chevron Research and Technology Company, unpublished

report.

12.03.2001

**Test condition** 

# OTHER RELEVANT INFORMATION (CAS NO.: 151006-62-1)

Test substance 1-dodecene trimer, hydrogenated (Alkane 4)

CAS 151006-62-1

**Type** other: The Potential Absorption and Metabolism of Alkane 4 (C36

Polyalphaolefin hydrogenated C12 trimer)

The aim of this report is to examine the literature data concerning the absorption and metabolism of Alkane 4 (a paraffin, a long chain (C36) branched alkane) and related substances in order to determine if there is a need to undertake toxicological testing at European Union level 1 and 2.

Normally, oral administration is the preferred route of administration for level 1 and 2 toxicity tests, thus this report concentrates on information on absorption from the gastro-intestinal tract. The limited information on the absorption of Alkane 4 was available from the base set tests. Other relevant information was needed. Data was also reviewed from shorter chain alkanes (C16-C20, C29 and C30), other polyalphaolefins, mineral oils, petroleum waxes and hydraulic fluids. Relevant tests are normally conducted in rats, therefore, the review concentrated on that species.

## Conclusion

The conclusions that can be drawn from the information in this review are:

- (1) Alkane 4 meets the US specification for mineral oils that may be used as components of non-food articles intended for use in contact with food and is a polyalphaolefin. It is a 36 carbon molecule.
- (2) The physicochemical parameters for Alkane 4 and the results of toxicity studies conducted as part of the base set test requirements suggest that Alkane 4 may not be absorbed.
- (3) Studies in the rat suggest that only limited amounts of the alkanes present in mineral hydrocarbons are absorbed, and that higher molecular weight material is less likely to be absorbed than lower molecular weight material.
- (4) Structure activity relationship studies indicate that alkanes with 36 carbon atoms are unlikely to be absorbed.
- (5) By analogy with C16-C29 alkanes, in the event of absorption taking place the first steps in any expected metabolism would be oxidation to the alcohol and formation fo the fatty acid. The product would enter the metabolic pathways described for fatty acids derived from food.

(6) Should absorption occur, it would be expected that the pathologic effects seen would be those associated with an excess intake of lipid like material. Although such effects were seen in rats following oral administration of mineral hydrocarbons, no pathologic effects were noted in studies whem mineral oils and waxes consisting of hydrocarbons with high molecular weights were fed.

In view of the need to pay proper attention to animal welfare and to minimize the use of animals in toxicity testing, these conclusions indicate that it would be inappropriate to undertake further toxicity studies on Alkane 4 at European Union levels 1 and 2.

Reliability

(1) valid without restriction

Reference

Illing, P. (2000) On the Potential Absorption and Metabolism of Alkane 4. Unpublished report prepasred for Chevron Chemical Company by Paul Illing Consultancy Services, Wirral, UK, and submitted to the UK Health and Safety Executive for Level 2 New Chemical Notification of Alkane 4.

# WATER SOLUBILITY (CAS No.: 68649-12-7)

Test Substance:	1-Decene tetramer/trimer; CAS 68649-12-7
Method/Guideline:	Calculated values using WSKOWWIN version 1.36, a subroutine of the computer program EPIWIN version 3.04
Year (guideline):	1999
Type (test type):	Not applicable
GLP:	Not applicable
Year (study performed):	Not applicable
Estimation Temperature:	25°C
Note: Concentration prep., vessel type, replication, test conditions.	Log Kow (octanol/water partition coefficient) values and a chemical structure are needed to calculate water solubility using the EPIWIN model. Water Solubility estimations performed by WSKOWWIN are based on a Kow correlation method described by W. Meylan, P. Howard and R. Boethling in "Improved method for estimating water solubility from octanol/water partition coefficient". <i>Environ. Toxicol. Chem.</i> <b>15</b> :100-106. 1995.  A C30 oligomer was used as a surrogate structure for 1-Decene tetramer/trimer.  SMILES: C=C(C=C(C=CCCCCCCCCCCCCCCCCCCCCCCCCCCC
Results: Units/Value:  Note: Deviations from	Calculated Chemical log K <sub>ow</sub> 1-Decene tetramer/trimer 14.62
protocol or guideline, analytical method.	Calculated WS (mg/L)  1-Decene tetramer/trimer  2.3 E <sup>-10</sup>
Conclusion:	Based on the calculated $K_{ow}$ value, decene tetramer/trimer can have a water solubility of < 1 ppb.
Reliability:	(2) Reliable with restrictions
	The value is calculated and represents a potential water solubility for decene tetramer/trimer.

Reference:	Meylan, M., SRC 1994-1999. WSKOWWIN is contained in the computer program EPIWIN. 1999. Estimation Program Interface for Windows, version 3.04. Syracuse Research Corporation, Syracuse, NY, USA.
Other (source):	ExxonMobil Biomedical Sciences, Inc.